CORRECTED VERSION

(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 21 March 2002 (21.03.2002)

PCT

(10) International Publication Number WO 02/022080 A3

(51) International Patent Classification7: C12N 15/86

(21) International Application Number: PCT/US01/28861

(22) International Filing Date:

14 September 2001 (14.09.2001)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

60/233,180

15 September 2000 (15.09.2000) US

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- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

- with international search report
- (88) Date of publication of the international search report:

 2 May 2002

 Date of publication of the revised international search report:

 16 January 2003
- (48) Date of publication of this corrected version:

6 March 2003

(15) Information about Corrections:

see PCT Gazette No. 10/2003 of 6 March 2003, Section II Previous Corrections:

see PCT Gazette No. 03/2003 of 16 January 2003, Section $\rm II$

see PCT Gazette No. 30/2002 of 25 July 2002, Section II

[Continued on next page]

(54) Title: ENHANCED FIRST GENERATION ADENOVIRUS VACCINES EXPRESSING CODON OPTIMIZED HIV1-GAG, POL, NEF AND MODIFICATIONS

(57) Abstract: First generation adenoviral vectors and associated recombinant adenovirus-based HIV vaccines which show enhanced stability and growth properties and greater cellular-mediated immunity are described within this specification. These adenoviral vectors are utilized to generate and produce through cell culture various adenoviral-based HIV-1 vaccines which contain HIV-1 gag, HIV-1 pol and/or HIV-1 nef polynucleotide pharmaceutical products, and biologically relevant modifications thereof. These adenovirus vaccines, when directly introduced into living vertebrate tissue, preferably a mammalian host such as a human or a non-human mammal of commercial or domestic veterinary importance, express the HIV1- Gag, Pol and/or Nef protein or biologically modification thereof, inducing a cellular immune response which specifically recognizes HIV-1. The exemplified polynucleotides of the present invention are synthetic DNA molecules encoding HIV-1 Gag, encoding codon optimized HIV-1 Pol, derivatives of optimized HIV-1 Pol (including constructs wherein protease, reverse transcriptase, RNAse H and integrase activity of HIV-1 Pol is inactivated), HIV-1 Nef and derivatives of optimized HIV-1 Nef, including nef mutants which effect wild type characteristics of Nef, such as myristylation and down regulation of host CD4. The adenoviral vaccines of the present invention, when administered alone or in a combined modality regime, will offer a prophylactic advantage to previously uninfected individuals and/or provide a therapeutic effect by reducing viral load levels within an infected individual, thus prolonging the asymptomatic phase of HIV-1 HIV

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

TITLE OF THE INVENTION ENHANCED FIRST GENERATION ADENOVIRUS VACCINES EXPRESSING CODON OPTIMIZED HIV1-GAG, POL, NEF AND MODIFICATIONS

5 CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims the benefit, under 35 U.S.C. §119(e), of U.S. provisional applications 60/233,180, 60/279,056, and Attorney Docket 20867PV2 (serial number unassigned), filed September 15, 2000, March 27, 2001, and September 7, 2001, respectively.

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STATEMENT REGARDING FEDERALLY-SPONSORED R&D Not Applicable

REFERENCE TO MICROFICHE APPENDIX

15 Not Applicable

FIELD OF THE INVENTION

The present invention relates to recombinant, replication-deficient first generation adenovirus vaccines found to exhibit enhanced growth properties and greater cellular-mediated immunity as compared to other replication-deficient vectors. The invention also relates to the associated first generation adenoviral vectors described herein, which, through the incorporation of additional 5' adenovirus sequence, enhance large scale production efficiency of the recombinant, replicationdefective adenovirus described herein. Another aspect of the instant invention is the surprising discovery that the intron A portion of the human cytomegalovirus (hCMV) promoter constitutes a region of instability in adenoviral vector constructs. Removal of this region from adenoviral expression constructs results in greatly improved vector stability. Therefore, improved vectors expressing a transgene under the control of an intron A-deleted CMV promoter constitute a further aspect of this invention. These adenoviral vectors are useful for generating recombinant adenovirus vaccines against human immunodeficiency virus (HIV). In particular, the first generation adenovirus vectors disclosed herein are utilized to construct and generate adenovirus-based HIV-1 vaccines which contain HIV-1 Gag, HIV-1 Pol and/or HIV-1 Nef polynucleotide pharmaceutical products, and biologically active modifications thereof. Host administration of the recombinant, replication-deficient adenovirus vaccines described herein results in expression of HIV-1 Gag, HIV-1- Pol and/or Nef protein or

immunologically relevant modifications thereof, inducing a cellular immune response which specifically recognizes HIV-1. The exemplified polynucleotides of the present invention are synthetic DNA molecules encoding codon optimized HIV-1 Gag, HIV-1 Pol, derivatives of optimized HIV-1 Pol (including constructs wherein protease, reverse transcriptase, RNAse H and integrase activity of HIV-1 Pol is inactivated), HIV-1 Nef, and derivatives of optimized HIV-1 Nef, including nef mutants which effect wild type characteristics of Nef, such as myristylation and down regulation of host CD4. The HIV adenovirus vaccines of the present invention, when administered alone or in a combined modality and/or prime/boost regimen, will offer a prophylactic advantage to previously uninfected individuals and/or provide a therapeutic effect by reducing viral load levels within an infected individual, thus prolonging the asymptomatic phase of HIV-1 infection.

BACKGROUND OF THE INVENTION

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Human Immunodeficiency Virus-1 (HIV-1) is the etiological agent of acquired human immune deficiency syndrome (AIDS) and related disorders. HIV-1 is an RNA virus of the Retroviridae family and exhibits the 5'LTR-gag-pol-env-LTR 3' organization of all retroviruses. The integrated form of HIV-1, known as the provirus, is approximately 9.8 Kb in length. Each end of the viral genome contains flanking sequences known as long terminal repeats (LTRs). The HIV genes encode at least nine proteins and are divided into three classes; the major structural proteins (Gag, Pol, and Env), the regulatory proteins (Tat and Rev); and the accessory proteins (Vpu, Vpr, Vif and Nef).

The gag gene encodes a 55-kilodalton (kDa) precursor protein (p55) which is expressed from the unspliced viral mRNA and is proteolytically processed by the HIV protease, a product of the pol gene. The mature p55 protein products are p17 (matrix), p24 (capsid), p9 (nucleocapsid) and p6.

The pol gene encodes proteins necessary for virus replication; a reverse transcriptase, a protease, integrase and RNAse H. These viral proteins are expressed as a Gag-Pol fusion protein, a 160 kDa precursor protein which is generated via a ribosomal frame shifting. The viral encoded protease proteolytically cleaves the Pol polypeptide away from the Gag-Pol fusion and further cleaves the Pol polypeptide to the mature proteins which provide protease (Pro, P10), reverse transcriptase (RT, P50), integrase (IN, p31) and RNAse H (RNAse, p15) activities.

The *nef* gene encodes an early accessory HIV protein (Nef) which has been shown to possess several activities such as down regulating CD4 expression, disturbing T-cell activation and stimulating HIV infectivity.

The *env* gene encodes the viral envelope glycoprotein that is translated as a 160-kilodalton (kDa) precursor (gp160) and then cleaved by a cellular protease to yield the external 120-kDa envelope glycoprotein (gp120) and the transmembrane 41-kDa envelope glycoprotein (gp41). Gp120 and gp41 remain associated and are displayed on the viral particles and the surface of HIV-infected cells.

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The tat gene encodes a long form and a short form of the Tat protein, a RNA binding protein which is a transcriptional transactivator essential for HIV-1 replication.

The rev gene encodes the 13 kDa Rev protein, a RNA binding protein. The Rev protein binds to a region of the viral RNA termed the Rev response element (RRE). The Rev protein promotes transfer of unspliced viral RNA from the nucleus to the cytoplasm. The Rev protein is required for HIV late gene expression and in turn, HIV replication.

Gp120 binds to the CD4/chemokine receptor present on the surface of helper T-lymphocytes, macrophages and other target cells in addition to other co-receptor molecules. X4 (macrophage tropic) virus show tropism for CD4/CXCR4 complexes while a R5 (T-cell line tropic) virus interacts with a CD4/CCR5 receptor complex. After gp120 binds to CD4, gp41 mediates the fusion event responsible for virus entry. The virus fuses with and enters the target cell, followed by reverse transcription of its single stranded RNA genome into the double-stranded DNA via a RNA dependent DNA polymerase. The viral DNA, known as provirus, enters the cell nucleus, where the viral DNA directs the production of new viral RNA within the nucleus, expression of early and late HIV viral proteins, and subsequently the production and cellular release of new virus particles. Recent advances in the ability to detect viral load within the host shows that the primary infection results in an extremely high generation and tissue distribution of the virus, followed by a steady state level of virus (albeit through a continual viral production and turnover during this phase), leading ultimately to another burst of virus load which leads to the onset of clinical AIDS. Productively infected cells have a half life of several days, whereas chronically or latently infected cells have a 3-week half life, followed by non-productively infected cells which have a long half life (over 100 days) but do not significantly contribute to day to day viral loads seen throughout the course of disease.

Destruction of CD4 helper T lymphocytes, which are critical to immune defense, is a major cause of the progressive immune dysfunction that is the hallmark of HIV infection. The loss of CD4 T-cells seriously impairs the body's ability to fight most invaders, but it has a particularly severe impact on the defenses against viruses, fungi, parasites and certain bacteria, including mycobacteria.

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Effective treatment regimens for HIV-1 infected individuals have become available recently. However, these drugs will not have a significant impact on the disease in many parts of the world and they will have a minimal impact in halting the spread of infection within the human population. As is true of many other infectious diseases, a significant epidemiologic impact on the spread of HIV-1 infection will only occur subsequent to the development and introduction of an effective vaccine. There are a number of factors that have contributed to the lack of successful vaccine development to date. As noted above, it is now apparent that in a chronically infected person there exists constant virus production in spite of the presence of anti-HIV-1 humoral and cellular immune responses and destruction of virally infected cells. As in the case of other infectious diseases, the outcome of disease is the result of a balance between the kinetics and the magnitude of the immune response and the pathogen replicative rate and accessibility to the immune response. Pre-existing immunity may be more successful with an acute infection than an evolving immune response can be with an established infection. A second factor is the considerable genetic variability of the virus. Although anti-HIV-1 antibodies exist that can neutralize HIV-1 infectivity in cell culture, these antibodies are generally virus isolate-specific in their activity. It has proven impossible to define serological groupings of HIV-1 using traditional methods. Rather, the virus seems to define a serological "continuum" so that individual neutralizing antibody responses, at best, are effective against only a handful of viral variants. Given this latter observation, it would be useful to identify immunogens and related delivery technologies that are likely to elicit anti-HIV-1 cellular immune responses. It is known that in order to generate CTL responses antigen must be synthesized within or introduced into cells, subsequently processed into small peptides by the proteasome complex, and translocated into the endoplasmic reticulum/Golgi complex secretory pathway for eventual association with major histocompatibility complex (MHC) class I proteins. CD8⁺ T lymphocytes recognize antigen in association with class I MHC via the T cell receptor (TCR) and the CD8 cell surface protein. Activation of naive CD8⁺ T cells into activated effector or memory cells generally requires both TCR engagement of antigen as described above as well as engagement of costimulatory proteins. Optimal

induction of CTL responses usually requires "help" in the form of cytokines from CD4⁺ T lymphocytes which recognize antigen associated with MHC class II molecules via TCR and CD4 engagement.

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European Patent Applications 0 638 316 (Published February 15, 1995) and 0 586 076 (Published March 9, 1994), (both assigned to American Home Products Corporation) describe replicating adenovirus vectors carrying an HIV gene, including env or gag. Various treatment regimens were used with chimpanzees and dogs, some of which included booster adenovirus or protein plus alum treatments.

Replication-defective adenoviral vectors harboring deletions in the E1 region are known, and recent adenoviral vectors have incorporated the known packaging repeats into these vectors; e.g., see EP 0 707 071, disclosing, *inter alia*, an adenoviral vector deleted of E1 sequences from base pairs 459 to 3328; and U.S. Patent No. 6,033,908, disclosing, *inter alia*, an adenoviral vector deleted of base pairs 459-3510. The packaging efficiency of adenovirus has been taught to depend on the number of incorporated individual A (packaging) repeats; *see*, *e.g.*, Gräble and Hearing, 1990 J. Virol. 64(5):2047-2056; Gräble and Hearing, 1992 J. Virol. 66(2):723-731.

Larder, et al., (1987, Nature 327: 716-717) and Larder, et al., (1989, Proc. Natl. Acad. Sci. 86: 4803-4807) disclose site specific mutagenesis of HIV-1 RT and the effect such changes have on *in vitro* activity and infectivity related to interaction with known inhibitors of RT.

Davies, et al. (1991, *Science* 252:, 88-95) disclose the crystal structure of the RNase H domain of HIV-1 Pol.

Schatz, et al. (1989, FEBS Lett. 257: 311-314) disclose that mutations Glu478Gln and His539Phe in a complete HIV-1 RT/RNase H DNA fragment results in defective RNase activity without effecting RT activity.

Mizrahi, et al. (1990, *Nucl. Acids. Res.* 18: pp. 5359-5353) disclose additional mutations Asp443Asn and Asp498Asn in the RNase region of the *pol* gene which also results in defective RNase activity. The authors note that the Asp498Asn mutant was difficult to characterize due to instability of this mutant protein.

Leavitt, et al. (1993, J. Biol. Chem. 268: 2113-2119) disclose several mutations, including a Asp64Val mutation, which show differing effect on HIV-1 integrase (IN) activity.

Wiskerchen, et al. (1995, J. Virol. 69: 376-386) disclose singe and double mutants, including mutation of aspartic acid residues which effect HIV-1 IN and viral replication functions.

It would be of great import in the battle against AIDS to produce a prophylactic- and/or therapeutic-based HIV vaccine which generates a strong cellular immune response against an HIV infection. The present invention addresses and meets these needs by disclosing a class of adenovirus vaccines which, upon host administration, express codon optimized and modified versions of the HIV-1 genes, gag, pol and nef. These recombinant, replication-defective adenovirus vaccines may be administered to a host, such as a human, alone or as part of a combined modality regimen and/or prime-boost vaccination regimen with components of the present invention and/or a distinct viral HIV DNA vaccine, non-viral HIV DNA vaccine, HIV subunit vaccine, an HIV whole killed vaccine and/or a live attenuated HIV vaccine.

SUMMARY OF THE INVENTION

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The present invention relates to enhanced replication-defective recombinant adenovirus vaccine vectors and associated recombinant, replication-deficient adenovirus vaccines which encode various forms of HIV-1 Gag, HIV-1 Pol, and/or HIV-1 Nef, including immunologically relevant modifications of HIV-1 Gag, HIV-1 Pol and HIV-1 Nef. The adenovirus vaccines of the present invention express HIV antigens and provide for improved cellular-mediated immune responses upon host administration. Potential vaccinees include but are not limited to primates and especially humans and non-human primates, and also include any non-human mammal of commercial or domestic veterinary importance. An effect of the improved recombinant adenovirus-based vaccines of the present invention should be a lower transmission rate to previously uninfected individuals (i.e., prophylactic applications) and/or reduction in the levels of the viral loads within an infected individual (i.e., therapeutic applications), so as to prolong the asymptomatic phase of HIV-1 infection. In particular, the present invention relates to adenoviral-based vaccines which encode various forms of codon optimized HIV-1 Gag (including but in no way limited to p55 versions of codon optimized full length (FL) Gag and tPA-Gag fusion proteins), HIV-1 Pol, HIV-1 Nef, and selected modifications of immunological relevance. The administration, intracellular delivery and expression of these adenovirus vaccines elicit a host CTL and Th response. The preferred replication-defective recombinant adenoviral vaccine vectors include but are not limited to synthetic DNA molecules which (1) encode codon optimized versions of wild type HIV-1 Gag; (2) encode codon optimized versions of HIV-1 Pol; (3) encode codon optimized versions of HIV-1 Pol fusion proteins; (4) encode codon optimized versions of modified HIV-1 Pol proteins and fusion proteins, including but not limited

to pol modifications involving residues within the catalytic regions responsible for RT, RNase and IN activity within the host cell; (5) encode codon optimized versions of wild type HIV-1 Nef; (6) codon optimized versions of HIV-1 Nef fusion proteins; and/or (7) codon optimized versions of HIV-1 Nef derivatives, including but not limited to nef modifications involving introduction of an amino-terminal leader sequence, removal of an amino-terminal myristylation site and/or introduction of dileucine motif mutations. The Nef-based fusion and modified proteins, disclosed within this specification and expressed from an adenoviral-based vector vaccine this specification, may possess altered trafficking and/or host cell function while retaining the ability to be properly presented to the host MHC I complex and in turn elicit a host CTL and Th response. Examples of HIV-1 Gag, Pol and/or Nef fusion proteins include but are not limited to fusion of a leader or signal peptide at the NH₂-teriminal portion of the viral antigen coding region. Such a leader peptide includes but is not limited to a tPA leader peptide.

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The adenoviral vector utilized in construction of the HIV-1 Gag-, HIV-1 Poland/or HIV-1 Nef- based vaccines of the present invention may comprise any replication-defective adenoviral vector which provides for enhanced genetic stability of the recombinant adenoviral genome through large scale production and purification of the recombinant virus. In other words, an HIV-1 Gag-, Pol- or Nef-based adenovirus vaccine of the present invention is a purified recombinant, replicationdefective adenovirus which is shown to be genetically stable through multiple passages in cell culture and remains so during large scale production and purification procedures. Such a recombinant adenovirus vector and harvested adenovirus vaccine lends itself to large scale dose filling and subsequent worldwide distribution procedures which will be demanded of an efficacious monovalent or multivalent HIV vaccine. The present invention meets this basic requirement with description of a replication-defective adenoviral vector and vectors derived therefrom, at least partially deleted in E1, comprising a wildtype adenovirus cis-acting packaging region from about base pair 1 to between from about base pair 342 (more preferably, 400) to about base pair 458 of the wildtype adenovirus genome. A preferred embodiment of the instant invention comprises base pairs 1-450 of a wildtype adenovirus. In other preferred embodiments, the replication -defective adenoviral vector has, in addition thereto, a region 3' to the E1-deleted region comprising base pairs 3511-3523. Basepairs 342-450 (more particularly, 400-450) constitute an extension of the 5'region of previously disclosed vectors carrying viral antigens, particularly HIV antigens (see, e.g., PCT International Application PCT/US00/18332, published

January 11, 2001 (WO 01/02067), which claims priority to U.S. Provisional Application Serial Nos. 60/142,631 and 60/148,981, filed 7/6/1999 and 8/13/1999, respectively; these documents herein incorporated by reference. Applicants have found that extending the 5' region further into the E1 gene into the disclosed vaccine vectors incorporated elements found to be important in optimizing the packaging of the virus.

As compared to previous vectors not comprising basepairs from about 1 to between from about base pair 342 (more preferably, 400) to about base pair 458 of the wildtype adenovirus genome, vectors comprising the above region exhibited enhanced growth characteristics, with approximately 5-10 fold greater amplification rates, a more potent virus effect, allowing lower doses of virus to be used to generate equivalent immunity; and a greater cellular-mediated immune response than replication-deficient vectors not comprising this region (basepairs 1-450). Even more important, adenoviral constructs derived therefrom are very stable genetically in large-scale production, particularly those comprising an expression cassette under the control of a hCMV promoter devoid of intron A. This is because Applicants have surprisingly found that the intron A portion of the hCMV promoter constituted a region of instability when employed in adenoviral vectors. Applicants have, therefore, identified an enhanced adenoviral vector which is particularly suited for use in gene therapy and nucleotide-based vaccine-vectors which, favorably, lends itself to large scale propagation.

A preferred embodiment of this invention is a replication-defective adenoviral vector in accordance with the above description wherein the gene is inserted in the form of a gene expression cassette comprising (a) a nucleic acid encoding a protein or biologically active and/or immunologically relevant portion thereof; (b) a heterologous promoter operatively linked to the nucleic acid of part a); and, (c) a transcription terminator.

In preferred embodiments, the E1 gene, other than that contained within basepairs 1-450 or, alternatively, that contained within base pairs 1-450 and 3511-3523 has been deleted from the adenoviral vector, and the gene expression cassette has replaced the deleted E1 gene. In other preferred embodiments, the replication defective adenovirus genome does not have a functional E3 gene, or the E3 gene has been deleted. Most preferably, the E3 region is present within the adenoviral genome. Further preferred embodiments are wherein the gene expression cassette is in an E1 anti-parallel (transcribed in a 3' to 5' direction relative to the vector backbone)

orientation or, more preferably, an E1 parallel (transcribed in a 5' to 3' direction relative to the vector backbone) orientation.

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Further embodiments relate to a shuttle plasmid vector comprising: an adenoviral portion and a plasmid portion, wherein said adenovirus portion comprises: a) a replication defective adenovirus genome, at least partially deleted in E1, comprising a wildtype adenovirus cis-acting packaging region from about base pair 1 to between from about base pair 342 (more preferably, 400) to about base pair 458 (preferably, 1-450) of the wildtype adenovirus genome and, preferably, in addition thereto, basepairs 3511-3523 of a wildtype adenovirus sequence; and b) a gene expression cassette comprising: (a) a nucleic acid encoding a protein or biologically active and/or immunologically relevant portion thereof; (b) a heterologous promoter operatively linked to the nucleic acid of part a); and (c) a transcription terminator and/or a polyadenylation site.

Other aspects of this invention include a host cell comprising said adenoviral vectors and/or said shuttle plasmid vectors; vaccine compositions comprising said vectors; and methods of producing the vectors comprising (a) introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and (b) harvesting the resultant adenoviral vectors.

To this end, the present invention particularly relates to harvested recombinant, replication defective virus derived from a host cell, such as but not limited to 293 cells or PER.C6® cells, including but not limited to harvested virus related to any of the MRKAd5 vector backbones, with or without an accompanying transgene, including but not limited to the HIV-1 antigens described herein. An HIV-1 vaccine is represented by any harvested, recombinant adenovirus material which expresses any one or more of the HIV-1 antigens disclosed herein. This harvested material may then be purified, formulated and stored prior to host administration.

Another aspect of this invention is a method of generating a cellular immune response against a protein in an individual comprising administering to the individual an adenovirus vaccine vector comprising:

a) a recombinant, replication defective adenoviral vector, at least partially deleted in E1, comprising a wildtype adenovirus *cis*-acting adenovirus packaging region from about base pair 1 to between from about base pair 342 (more preferably, 400) to about base pair 458 (preferably, 1-450) and, preferably in addition thereto, base pairs 3511-3523 of a wildtype adenovirus sequence, and,

b) a gene expression cassette comprising:(i) a nucleic acid encoding a protein or biologically active and/or immunologically relevant portion thereof; (ii) a heterologous promoter operatively linked to the nucleic acid of part a); and (iii) a transcription terminator and/or a polyadenylation site.

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In view of the efficacious nature of the adenoviral and/or DNA plasmid vaccines described herein, the present invention relates to all methodology regarding administration of one or more of these adenoviral and/or DNA plasmid vaccines to provide effective immunoprophylaxis, to prevent establishment of an HIV-1 infection following exposure to this virus, or as a post-HIV infection therapeutic vaccine to mitigate the acute HIV-1 infection so as to result in the establishment of a lower virus load with beneficial long term consequences. As discussed herein, such a treatment regimen may include a monovalent or multivalent composition, various combined modality applications, and/or a prime/boost regimen to as to optimize antigen expression and a concomitant cellular-mediated and/or humoral immune response upon inoculation into a living vertebrate tissue. Therefore, the present invention provides for methods of using the adenoviral and/or DNA plasmid vaccines disclosed herein within the various parameters disclosed herein as well as any additional parameters known in the art, which, upon introduction into mammalian tissue induces intracellular expression of the gag, pol and/or nef-based vaccines.

To this end, the present invention relates in part to methods of generating a cellular immune response in a vaccinee, preferably a human vaccinee, wherein the individual is given more than one administration of adenovirus vaccine vector, and it may be given in a regimen accompanied by the administration of a plasmid vaccine. The plasmid vaccine (also referred to herein as a "DNA plasmid vaccine" or "vaccine plasmid" comprises a nucleic acid encoding a protein or an immunologically relevant portion thereof, a heterologous promoter operably linked to the nucleic acid sequence, and a transcription terminator or a polyadenylation signal (such as bGH or SPA, respectively). There may be a predetermined minimum amount of time separating the administrations. The individual can be given a first dose of plasmid vaccine, and then a second dose of plasmid vaccine. Alternatively, the individual may be given a first dose of adenovirus vaccine, and then a second dose of adenovirus vaccine. In other embodiments, the plasmid vaccine is administered first, followed after a time by administration of the adenovirus vaccine. Conversely, the adenovirus vaccine may be administered first, followed by administration of plasmid vaccine after a time. In these embodiments, an individual may be given multiple doses of the same adenovirus serotype in either viral vector or plasmid form, or the virus may be of

differing serotypes. In the alternative, a viral antigen of interest can be first delivered via a viral vaccine other than an adenovirus-based vaccine, and then followed with the adenoviral vaccine disclosed. Alternative viral vaccines include but are not limited to pox virus and venezuelan equine encephilitis virus.

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The present invention also relates to multivalent adenovirus vaccine compositions which comprise Gag, Pol and Nef components described herein; see, e.g., Example 29 and Table 25. Such compositions will provide for an enhanced cellular immune response subsequent to host administration, particularly given the genetic diversity of human MHCs and of circulating virus. Examples, but not limitations, include MRKAd5-vector based multivalent vaccine compositions which provide for a divalent (i.e., gag and nef, gag and pol, or pol and nef components) or a trivalent vaccine (i.e., gag, pol and nef components) composition. Such a mutlivalent vaccine may be filled for a single dose or may consist of multiple inoculations of each individually filled component; and may in addition be part of a prime/boost regimen with viral or non-viral vector vaccines as introduced in the previous paragraph. To this end, preferred compositions are MRKAd5 adenovirus used in combination with multiple, distinct HIV antigen classes. Each HIV antigen class is subject to sequence manipulation, thus providing for a multitude of potential vaccine combinations; and such combinations are within the scope of the present invention. The utilization of such combined modalities vaccine formulation and administration increase the probability of eliciting an even more potent cellular immune response when compared to inoculation with a single modality regimen.

The concept of a "combined modality" as disclosed herein also covers the alternative mode of administration whereby multiple HIV-1 viral antigens may be ligated into a proper shuttle plasmid for generation of a pre-adenoviral plasmid comprising multiple open reading frames. For example, a trivalent vector may comprise a gag-pol-nef fusion, in either a E3(-) or E3(+) background, preferably a E3 deleted backbone, or possibly a "2+1" divalent vaccine, such as a gag-pol fusion (i.e., codon optimized p55 gag and inactivated optimized pol; Example 29 and Table 25) within the same MRKAd5 backbone, with each open reading frame being operatively linked to a distinct promoter and transcription termination sequence. Alternatively, the two open reading frames may be operatively linked to a single promoter, with the open reading frames operatively linked by an internal ribosome entry sequence (IRES). Therefore, a multivalent vaccine delivered as a single, or possibly a second harvested recombinant, replication-deficient adenovirus is contemplated as part of the present invention.

Therefore, the adenoviral vaccines and plasmid DNA vaccines of this invention may be administered alone, or may be part of a prime and boost administration regimen. A mixed modality priming and booster inoculation scheme will result in an enhanced immune response, particularly if pre-existing anti-vector immune responses are present. This one aspect of this invention is a method of priming a subject with the plasmid vaccine by administering the plasmid vaccine at least one time, allowing a predetermined length of time to pass, and then boosting by administering the adenoviral vaccine. Multiple primings typically, 1-4, are usually employed, although more may be used. The length of time between priming and boost may typically vary from about four months to a year, but other time frames may be used. In experiments with rhesus monkeys, the animals were primed four times with plasmid vaccines, then were boosted 4 months later with the adenoviral vaccine. Their cellular immune response was notably higher than that of animals which had only received adenoviral vaccine. The use of a priming regimen may be particularly preferred in situations where a person has a pre-existing anti-adenovirus immune response.

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It is an object of the present invention to provide for enhanced replication-defective recombinant adenoviral vaccine vector backbones. These recombinant adenoviral backbones may accept one or more transgenes, which may be passaged through cell culture for growth, amplification and harvest.

It is a further object to provide for enhanced replication-defective recombinant adenoviral vaccine vectors which encode various transgenes.

It is also an object of the present invention to provide for a harvested recombinant, replication-deficient adenovirus which shows enhanced growth and amplification rates while in combination with increased virus stability after continuous passage in cell culture. Such a recombinant adenovirus is particularly suited for use in gene therapy and nucleotide-based vaccine vectors which, favorably, lends itself to large scale propagation.

To this end, it is an object of the present invention to provide for (1) enhanced replication-defective recombinant adenoviral vaccine vectors as described herein which encode various forms of HIV-1 Gag, HIV-1 Pol, and/or HIV-1 Nef, including immunologically relevant modifications of HIV-1 Gag, HIV-1 Pol and HIV-1 Nef, and (2) harvested, purified recombinant replication-deficient adenovirus generated by passage of the adenoviral vectors of (1) through one or multiple passages through cell culture, including but not limited to passage through 293 cells or PER.C6[®] cells.

It is also an object of the present invention to provide for recombinant adenovirus harvested by one or multiple passages through cell culture. As relating to recombinant adenoviral vaccine vector, this recombinant virus is harvested and formulated for subsequent host administration.

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It is also an object of the present invention to provide for replication-defective adenoviral vectors wherein at least one gene is inserted in the form of a gene expression cassette comprising (a) a nucleic acid encoding a protein or biologically active and/or immunologically relevant portion thereof; (b) a heterologous promoter operatively linked to the nucleic acid of part a); and, (c) a transcription terminator.

10 It is also an object of the present invention to provide for a host cell comprising said adenoviral vectors and/or said shuttle plasmid vectors; vaccine compositions comprising said vectors; and methods of producing the vectors comprising (a) introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and (b) harvesting the resultant adenoviral vectors. It is a further object of the present invention to provide for methods of generating a 15 cellular immune response against a protein in an individual comprising administering to the individual an adenovirus vaccine vector comprising a) a replication defective adenoviral vector, at least partially deleted in E1, comprising a wildtype adenovirus cis-acting packaging region from about base pair 1 to between from about base pair 20 342 (more preferably, 400) to about 450 (preferably, 1-450) and, preferably, 3511-3523 of a wildtype adenovirus sequence, and, b) a gene expression cassette comprising:(i) a nucleic acid encoding a protein or biologically active and/or immunologically relevant portion thereof; (ii) a heterologous promoter operatively linked to the nucleic acid of part a); and (iii) a transcription terminator and/or a 25 polyadenylation site.

It is also an object of the present invention to provide various alternatives for vaccine administration regimes, namely administration of one or more adenoviral and/or DNA plasmid vaccines described herein to provide effective immunoprophylaxis for uninfected individuals or a therapeutic treatment for HIV infected patients. Such processes include but are not limited to multivalent HIV-1 vaccine compositions, various combined modality regimes as well as various prime/boost alternatives. These methods of administration, relating to vaccine composition and/or scheduled administration, will increase the probability of eliciting an even more potent cellular immune response when compared to inoculation with a single modality regimen.

As used throughout the specification and claims, the following definitions and abbreviations are used:

"HAART" refers to - highly active antiretroviral therapy -.

"first generation" vectors are characterized as being replication-defective.

5 They typically have a deleted or inactivated E1 gene region, and preferably have a deleted or inactivated E3 gene region as well.

"AEX" refers to Anion Exchange chromatography.

"QPA" refers to Quick PCR-based Potency Assay.

"bps" refers to basepairs.

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"s" or "str" denotes that the transgene is in the E1 parallel or "straight" orientation.

"PBMCs" refers to peripheral blood monocyte cells.

"FL" refers to full length.

"FLgag" refers to a full-length optimized gag gene, as shown in Figure 2.

"Ad5-Flgag" refers to an adenovirus serotype 5 replication deficient virus which carries an expression cassette which comprises a full length optimized gag gene under the control of a CMV promoter.

"Promoter" means a recognition site on a DNA strand to which an RNA polymerase binds. The promoter forms an initiation complex with RNA polymerase to initiate and drive transcriptional activity. The complex can be modified by activating sequences such as enhancers or inhibiting sequences such as silencers.

"Leader" means a DNA sequence at the 5' end of a structural gene which is transcribed along with the gene. This usually results a protein having an N-terminal peptide extension, often referred to as a pro-sequences.

"Intron" means a section of DNA occurring in the middle of a gene which does not code for an amino acid in the gene product. The precursor RNA of the intron is excised and is therefore not transcribed into mRNA not translated into protein.

"Immunologically relevant" or "biologically active" means (1) with regards to a viral protein, that the protein is capable, upon administration, of eliciting a measurable immune response within an individual sufficient to retard the propagation and/or spread of the virus and/or to reduce the viral load present within the individual; or (2) with regards to a nucleotide sequence, that the sequence is capable of encoding for a protein capable of the above.

"Cassette" refers to a nucleic acid sequence which is to be expressed, along with its transcription and translational control sequences. By changing the cassette, a vector can express a different sequence.

"bGHpA" refers to the bovine growth hormone transcription terminator/polyadenylation sequence.

"tPAgag" refers to a fusion between the leader sequence of the tissue plasminogen activator leader sequence and an optimized HIV gag gene, as exemplified in Figure 30A-B, whether in a DNA or adenovirus-based vaccine vector.

Where utilized, "IA" or "inact" refers to an <u>inactivated</u> version of a gene (e.g. IApol).

"MCS" is "multiple cloning site".

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In general, adenoviral constructs, gene constructs are named by reference to the genes contained therein. For example:

"Ad5 HIV-1 gag", also referred to as the original HIV-1 gag adenoviral vector, is a vector containing a transgene cassette composed of a hCMV intron A promoter, the full length version of the human codon-optimized HIV-1 gag gene, and the bovine growth hormone polyadenylation signal. The transgene was inserted in the E1 antiparallel orientation in an E1 and E3 deleted adenovector.

"MRK Ad5 HIV-1 gag" also referred to as "MRKAd5gag" or "Ad5gag2" is an adenoviral vector taught herein which is deleted of E1, comprises basepairs 1-450 and 3511-3523, and has a human codon-optimized HIV-1 gene in an E1 parallel orientation under the control of a CMV promoter without intron A. The construct also comprises a bovine growth hormone polyadenylation signal.

"pV1JnsHIVgag", also referred to as "HIVFLgagPR9901", is a plasmid comprising the CMV immediate-early (IE) promoter and intron A, a full-length codon-optimized HIV gag gene, a bovine growth hormone-derived polyadenylation and transcriptional termination sequence, and a minimal pUC backbone.

"pV1JnsCMV(no intron)-FLgag-bGHpA" is a plasmid derived from pV1JnsHIVgag which is deleted of the intron A portion of CMV and which comprises the full length HIV gag gene. This plasmid is also referred to as "pV1JnsHIVgag-bGHpA", pV1Jns-hCMV-FL-gag-bGHpA" and "pV1JnsCMV(no intron) + FLgag + bGHpA".

"pV1JnsCMV(no intron)-FLgag-SPA" is a plasmid of the same composition as pV1JnsCMV(no intron)-FLgag-bGHpA except that the SPA termination sequence replaces that of bGHpA. This plasmid is also referred to as "pV1Jns-HIVgag-SPA" and pV1Jns-hCMV-FLgag-SPA".

"pdelE1sp1A" is a universal shuttle vector with no expression cassette (i.e., no promoter or polyA). The vector comprises wildtype adenovirus serotype 5 (Ad5) sequences from bp 1 to bp 341 and bp 3524 to bp 5798, and has a multiple cloning

site between the Ad5 sequences ending 341 bp and beginning 3524 bp. This plasmid is also referred to as the original Ad 5 shuttle vector.

"MRKpdelE1sp1A" or "MRKpdelE1(Pac/pIX/pack450)" or

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"MRKpdelE1(Pac/pIX/pack450)Cla1" is a universal shuttle vector with no expression cassette (i.e. no promoter or polyA) comprising wildtype adenovirus serotype 5 (Ad5) sequences from bp1 to bp450 and bp 3511 to bp 5798. The vector has a multiple cloning site between the Ad5 sequence ending 450 bp and beginning 3511 bp. This shuttle vector may be used to insert the CMV promoter and the bGHpA fragments in both the straight ("str". or E1 parallel) orientation or in the opposite (opp. or E1 antiparallel) orientation)

"MRKpdelE1(Pac/pIX/pack450)+CMVmin+BGHpA(str.)" is still another shuttle vector which is the modified vector that contains the CMV promoter (no intronA) and the bGHpA fragments. The expression unit containing the hCMV promoter (no intron A) and the bovine growth hormone polyadenylation signal has been inserted into the shuttle vector such that insertion of the gene of choice at a unique BgIII site will ensure the direction of transcription of the transgene will be Ad5 E1 parallel when inserted into the MRKpAd5(E1/E3+)Cla1 pre-plasmid. This shuttle vector, as shown in Figures 22 and 23, was used to insert the respective IApol and G2A,LLAA nef genes directly into.

"MRKpdelE1-CMV(no intron)-FLgag-bGHpA" is a shuttle comprising Ad5 sequences from basepairs 1-450 and 3511-5798, with an expression cassette containing human CMV without intron A, the full-length human codon-optimized HIV gag gene and bovine growth hormone polyadenylation signal. This plasmid is also referred to as "MRKpdelE1 shuttle +hCMV-FL-gag-BGHpA"

"MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA" is an adenoviral vector comprising all Ad5 sequences except those nucleotides encompassing the E1 region (from 451-3510), a human CMV promoter without intron A, a full-length human codon-optimized HIV gag gene, and a bovine growth hormone polyadenylation signal. This vector is also referred to as "MRKpAdHVE3 + hCMV-FL-gag-BGHpA", "MRKpAd5HIV-1gag", "MRKpAd5gag", "pMRKAd5gag" or "pAd5gag2".

"pV1Jns-HIV-pol inact(opt)" or "pV1Jns-HIV IA pol (opt) is the inactivated Pol gene (contained within SEQ ID NO:3) cloned into the BgIII site of V1Jns (Figure 17A-C). As noted herein, various derivatives of HIV-1 pol may be cloned into a plasmid expression vector such as V1Jns or V1Jns-tPA, thus serving directly as DNA vaccine candidates or as a source for subcloning into an appropriate adenoviral vector.

"MRKpdel+hCMVmin+FL-pol+bGHpA(s)" is the "MRKpdelE1(Pac/pIX/pack450)+CMVmin+BGHpA(str.)" shuttle mentioned above which contains the IA pol gene is the proper orientation. This shuttle vector is used in a bacterial recombination with MRKpAd(E1-/E3+)Cla1.

"MRKpAd+hCMVmin+FL-pol+bGHpA(S)E3+", also referred to herein as "pMRKAd5pol", is the pre-adenovirus plasmid which comprises a CMV-pol inact(opt)-pGHpA construct. The construction of this pre-adenovirus plasmid is shown in Figure 22.

"pV1Jns/nef (G2A,LLAA)" or "V1Jns/opt nef (G2A,LLAA)" comprises codon optimized HIV-1 Nef wherein the open reading frame codes for modifications at the amino terminal myristylation site (Gly-2 to Ala-2) and substitution of the Leu-174-Leu-175 dileucine motif to Ala-174-Ala-175 (SEQ ID NO:13; which comprises an initiating methionine residue at nucleotides 12-14 and a "TAA" stop codon from nucleotides 660-662). This fragment is subcloned into the Bgl II site of V1Jns and/orV1Jns-tPA (Figures 16A-B). As noted above for HIV-1 pol, HIV-1 nef constructs may be cloned into a plasmid expression vector such as V1Jns or V1Jns-tPA, thus serving directly as DNA vaccine candidates or as a source for subcloning into an appropriate adenoviral vector.

"MRKpdelE1hCMVminFL-nefBGHpA(s)", also referred to herein as "pMRKAd5nef", is the pre-adenovirus plasmid which comprises a CMV-nef (G2A,LLAA) codon optimized sequence. The construction of this pre-adenovirus plasmid is shown in Figure 23.

BRIEF DESCRIPTION OF THE FIGURES

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Figure 1 shows the original HIV-1 gag adenovector (Ad5HIV-1gag). This vector is disclosed in PCT International Application No. PCT/US00/18332 (WO 01/02607) filed July 3, 2000, claiming priority to U.S. Provisional Application Serial No. 60/142,631, filed July 6, 1999 and U.S. Application Serial No. 60/148,981, filed August 13, 1999, all three applications which are hereby incorporated by reference.

Figure 2 shows the nucleic acid sequence (SEQ ID NO: 29) of the optimized human HIV-1 gag open reading frame.

Figure 3 shows diagrammatically the new transgene constructs in comparison with the original gag transgene.

Figure 4 shows the modifications made to the original adenovector backbone in the generation of the novel vectors of the instant invention.

Figure 5 shows the virus mixing experiments that were carried out to determine the effects of the addition made to the packaging signal region (Expt. #1) and the E3 gene on viral growth (Expt. #2). The bars denote the region of modifications made to the E1 deletion.

Figure 6 shows an autoradiograph of viral DNA analysis following the viral mixing experiments described in Examples 6 and 7.

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Figures 7A, 7B and 7C are as follows: Figure 7A shows the hCMV-Flgag-bGHpA adenovectors constructed within the MRKpAdHVE3 and MRKpAdHVO adenovector backbones. Both E1 parallel and E1 antiparallel transgene orientation are represented. Figure 7B shows the hCMV-Flgag-SPA adenovectors constructed within the MRKpAdHVE3 and MRKpAdHVO adenovector backbones. Again, both E1 parallel and E1 antiparallel transgene orientation are represented. Figure 7C shows the mCMV-Flgag-bGHpA adenovectors constructed within the MRKpAdHVE3 and MRKpAdHVO adenovector backbones. Once again, both E1 parallel and E1 antiparallel transgene orientation are represented.

Figure 8A shows the experiment designed to test the effect of transgene orientation.

Figure 8B shows the experiments designed to test the effect of polyadenylation signal.

Figure 9 shows viral DNA from the four adenoviral vectors tested (Example 12) at P5, following *Bst*E11 digestion.

Figure 10 shows viral DNA analysis of passages 11 and 12 of MRKpAdHVE3, MRKAd5HIV-1gag, and MRKAd5HIV-1gagE3-.

Figure 11 shows viral DNA analysis (*Hind*III digestion) of passage 6 MRKpAdHVE3 and MRKAd5HIV-1gag used to initiate the viral competition study. The last two lanes are passage 11 analysis of duplicate passages of the competition study (each virus at MOI of 280 viral particles).

Figure 12 shows viral DNA analysis by *Hind* III digestion on high passage numbers for MRKAd5HIV-1gag in serum-containing media with collections made at specified times. The first lane shows the 1kb DNA size marker. The other lanes represent pre-plasmid control (digested with Pac1 and *Hind*III), MRKAd5HIV-1gag at P16, P19, and P21.

Figure 13 shows serum anti-p24 levels at 3 wks post i.m. immunization of balb/c mice (n=10) with varying doses of several Adgag constructs: (A) MRK Ad5 HIV-1 gag (through passage 5); (B) MRKAd5 hCMV-FLgag-bGHpA (E3-); (C) MRKAd5 hCMV-FLgag-SPA (E3+); (D) MRKAd5 mCMV-FLgag-bGHpA (E3+);

(E) research lot (293 cell-derived) of Ad5HIV-1 gag; and (F) clinical lot (Ad5gagFN0001) of Ad5HIV-1 gag. Reported are the geometric mean titers (GMT) for each cohort along with the standard error bars.

Figure 14 shows a restriction map of the pMRKAd5HIV-1gag vector.

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Figures 15A-X illustrates the nucleotide sequence of the pMRKAd5HIV-1gag vector (SEO ID NO:27.[coding] and SEQ ID NO:28 [non-coding]).

Figures 16A-B shows a schematic representation of DNA vaccine expression vectors V1Jns (A) and V1Jns-tPA (B), which are utilized for HIV-1 gag, pol and nef constructs in various DNA/viral vector combined modality regimens as disclosed herein.

Figures 17A-C shows the nucleotide (SEQ ID NO:3) and amino acid sequence (SEQ ID NO:4) of IA-Pol. Underlined codons and amino acids denote mutations, as listed in Table 1.

Figure 18 shows codon optimized nucleotide and amino acid sequences through the fusion junction of tPA-pol inact(opt) (contained within SEQ ID NOs: 7 and 8, respectively). The underlined portion represents the NH₂-terminal region of IA-Pol.

Figures 19A-B show a nucleotide sequence comparison between wild type nef(jrfl) and codon optimized nef. The wild type nef gene from the jrfl isolate consists of 648 nucleotides capable of encoding a 216 amino acid polypeptide. WT, wild type sequence (SEQ ID NO:19); opt, codon-optimized sequence (contained within SEQ ID NO:1). The Nef amino acid sequence is shown in one-letter code (SEQ ID NO:2).

Figures 20A-C show nucleotide sequences at junctions between nef coding sequence and plasmid backbone of nef expression vectors V1Jns/nef (Figure 20A), V1Jns/nef(G2A,LLAA) (Figure 20B), V1Jns/tpanef (Figure 20C) and V1Jns/tpanef(LLAA) (Figure 20C, also). 5' and 3' flanking sequences of codon optimized nef or codon optimized nef mutant genes are indicated by bold/italic letters; nef and nef mutant coding sequences are indicated by plain letters. Also indicated (as underlined) are the restriction endonuclease sites involved in construction of respective nef expression vectors. V1Jns/tpanef and V1Jns/tpanef(LLAA) have identical sequences at the junctions.

Figure 21 shows a schematic presentation of nef and nef derivatives. Amino acid residues involved in Nef derivatives are presented. Glycine 2 and Leucine174 and 175 are the sites involved in myristylation and dileucine motif, respectively. For both versions of the tpanef fusion genes, the putative leader peptide cleavage sites are

indicated with "*", and a exogenous serine residue introduced during the construction of the mutants is underlined.

Figure 22 shows diagrammatically the construction of the pre-adenovirus plasmid construct, MRKAd5Pol.

Figure 23 shows diagrammatically the construction of the pre-adenovirus plasmid construct, MRKAd5Nef.

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Figure 24 shows a comparison of clade B vs. clade C anti-gag T cell responses in clade B HIV-infected subjects.

Figure 25 shows a comparison of clade B vs. clade C anti-nef T cell responses in clade B HIV-infected subjects.

Figures 26A-AO illustrates the nucleotide sequence of the pMRKAd5HIV-1pol adenoviral vector (SEQ ID NO:32 [coding] and SEQ ID NO:33 [non-coding]), comprising the coding region of the inactivated pol gene (SEQ ID NO3).

Figures 27A-AM illustrates the nucleotide sequence of the pMRKAd5HIV-1 nef adenoviral vector (SEQ ID NO:34 [coding] and SEQ ID NO:35 [non-coding]), comprising the coding region of the inactivated pol gene (SEQ ID NO13).

Figure 28 shows the stability of MRKAd5 vectors comprising various promoter fragments (hCMV or mCMV) and terminations signals (bGH or SPA) in E3(+) or E3(-) backbones.

Figures 29A and B shows the anion-exchange HPLC viral particle concentrations of the freeze-thaw recovered cell associated virus at the 24, 36, 48, and 60 hpi time points (Figure 29A) and the timcourse QPA supernatant titers (Figure 29B) for MRKAd5gag, MRKAd5pol and MRKAd5nef.

Figure 30 shows the nucleotide sequence (SEQ ID NO:36) and amino acid sequence (SEQ ID NO:37) comprising the open reading frame of a representative tPA-gag fusion for use in the DNA and/or adenoviral vaccine disclosed herein.

Figure 31 shows the intracellular γIFN staining of PBMCs collected at week 10 (post DNA prime) and week 30 (post Ad boost). The cells were stimulated overnight in the presence or absence of the gag peptide pool. They were subsequently stained using fluorescence-tagged anti-CD3, anti-CD8, anti-CD4, and anti-γIFN monoclonal antibodies. Each plot shows all CD3+ T cells which were segregated in terms of positive staining for surface CD8 and γIFN production. The numbers in the upper right and lower right quadrants of each plot are the percentages of CD3⁺ cells that were CD8⁺γIFN⁺ and CD4⁺γIFN⁺, respectively.

Figure 32 shows a comparison of single-modality adenovirus immunization with DNA + adjuvant prime/adenovirus boost immunization.

Figures 33A-B show the nucleotide sequence (SEQ ID NO: 38) of the open reading frame for the gag-IApol fusion of Example 29.

Figures 34A-B show the protein sequence (SEQ ID NO:39) of the gag-IApol fustion frame.

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DETAILED DESCRIPTION OF THE INVENTION

A novel replication-defective, or "first generation," adenoviral vector suitable for use in gene therapy or nucleotide-based vaccine vectors is described. This vector is at least partially deleted in E1 and comprises a wildtype adenovirus cis-acting packaging region from about base pair 1 to between about base pair 342 (more preferably, 400) to about 458 (preferably, 1-450) and, preferably, 3511-3523 of a wild-type adenovirus sequence. It has been found that a vector of this description possesses enhanced growth characteristics, with approximately 5-10 fold greater amplification rates, and is more potent allowing lower doses of virus to be used to generate equivalent immunity. The vector, furthermore, generates a harvested recombinant adenovirus which shows greater cellular-mediated immune responses than replication-deficient vectors not comprising this region (basepairs 342-450). Adenoviral constructs derived from these vectors are, further, very stable genetically, particularly those comprising a transgene under the control of a hCMV promoter devoid of intron A. Viruses in accordance with this description were passaged continually and analyzed; see Example 12. Each virus analyzed maintained it correct genetic structure. Analysis was also carried out under propagation conditions similar to that performed in large scale production. Again, the vectors were found to possess enhanced genetic stability; see Figure 12. Following 21 passages, the viral DNA showed no evidence of rearrangement, and was highly reproducible from one production lot to the next. The outcome of all relevant tests indicate that the adenoviral vector is extremely well suited for large-scale production of recombinant, replication-deficient adenovirus, as shown herein with the data associated with Figure 28.

A preferred adenoviral vector in accordance with this description is a vector comprising basepairs 1-450, which is deleted in E3. This vector can accommodate up to approximately 7,500 base pairs of foreign DNA inserts (or exogenous genetic material). Another preferred vector is one retaining E3 which comprises basepairs 1-450. A preferred vector of this description is an E3+ vector comprising basepairs 1-450 and 3511-3523. This vector, when deleted of the region spanning basepairs 451-3510, can accommodate up to approximately, 4,850 base pairs of foreign DNA inserts

(or exogenous genetic material). The cloning capacities of the above vectors have been determined using 105% of the wildtype Ad5 sequence as the upper genome size limit.

Wildtype adenovirus serotype 5 is used as the basis for the specific basepair numbers provided throughout the specification. The wildtype adenovirus serotype 5 sequence is known and described in the art; see, Chroboczek et al., 1992 J. Virology 186:280, which is hereby incorporated by reference. Accordingly, a particular embodiment of the instant invention is a vector based on the adenovirus serotype 5 sequence. One of skill in the art can readily identify the above regions in other adenovirus serotypes (e.g., serotypes 2, 4, 6, 12, 16, 17, 24, 31, 33, and 42), regions defined by basepairs corresponding to the above basepair positions given for adenovirus serotype 5. Accordingly, the instant invention encompasses all adenoviral vectors partially deleted in E1 comprising basepairs corresponding to 1-450 (particularly, 342-450) and, preferably, 3511-3523 of a wild-type adenovirus serotype 5 (Ad5) nucleic acid sequence. Particularly preferred embodiments of the instant invention are those derived from adenoviruses like Ad5 which are classified in subgroup C (e.g., Ad2).

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Vectors in accordance with the instant invention are at least partially deleted in E1. Preferably the E1 region is completely deleted or inactivated. Most preferably, the region deleted of E1 is within basepairs 451-3510. It is to be noted that the extended 5' and 3' regions of the disclosed vectors are believed to effectively reduce the size of the E1 deletion of previous constructs without overlapping any part of the E1A/E1B gene present in the cell line used, i.e., the PER.C6® cell line transefected with base pairs 459-3510. Overlap of adenoviral sequences is avoided because of the possibility of recombination. One of ordinary skill in the art can certainly appreciate that the instant invention can, therefore, be modified if a different cell line transfected with a different segment of adenovirus DNA is utilized. For purposes of exemplification, a 5' region of base pairs 1 to up to 449 is more appropriate if a cell line is transfected with adenoviral sequence from base pairs 450-3510. This holds true as well in the consideration of segments 3' to the E1 deletion.

Preferred embodiments of the instant invention possess an intact E3 region (i.e., an E3 gene capable of encoding a functional E3). Alternate embodiments have a partially deleted E3, an inactivated E3 region, or a sequence completely deleted of E3. Applicants have found, in accordance with the instant invention, that virus comprising the E3 gene were able to amplify more rapidly compared with virus not comprising an E3 gene; see Figure 6 wherein a diagnostic CsCl band corresponding to the E3+ virus

tested (5,665 bp) was present in greater amount compared with the diagnostic band of 3,010 bp corresponding to the E3- virus. These results were obtained following a virus competition study involving mixing equal MOI ratio (1:1) of adenovectors both comprising the E3 gene and not comprising the E3 gene. This increased amplification capacity of the E3+ adenovectors was subsequently confirmed with growth studies; see Table 4A, wherein the E3+ virus exhibit amplification ratios of 470, 420 and 320 as compared with the 115 and 40-50 of the E3- constructs.

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As stated above, vectors in accordance with the instant invention can accommodate up to approximately 4,850 base pairs of exogenous genetic material for an E3+ vector and approximately 7,500 base pairs for an E3- vector. Preferably, the insert brings the adenoviral vector as close as possible to a wild-type genomic size (e.g., for Ad5, 35,935 basepairs). It is well known that adenovirus amplifies best when they are close to their wild-type genomic size.

The genetic material can be inserted in an E1-parallel or an E1 anti-parallel orientation, as such is illustrated in Figure 7A, 7B, 7C and Figure 8A. Particularly preferred embodiments of the instant invention, have the insert in an E1-parallel orientation. Applicants have found, via competition experiments with plasmids containing transgenes in differing orientation (Figure 8A), that vector constructs with the foreign DNA insert in an E1-parallel orientation amplify better and actually outcompete E1-antiparallel-oriented transgenes. Viral DNA analysis of the mixtures at passage 3 and certainly at passage 6, showed a greater ratio of the virus carrying the transgene in the E1 parallel orientation as compared with the E1 anti-parallel version. By passage 10, the only viral species observed was the adenovector with the transgene in the E1 parallel orientation for both transgenes tested.

Adenoviral vectors in accordance with the instant invention are particularly well suited to effectuate expression of desired proteins, one example of which is an HIV protein, particularly an HIV full length gag protein. Exogenous genetic material encoding a protein of interest can exist in the form of an expression cassette. A gene expression cassette preferably comprises (a) a nucleic acid encoding a protein of interest; (b) a heterologous promoter operatively linked to the nucleic acid encoding the protein; and (c) a transcription terminator.

The transcriptional promoter is preferably recognized by an eukaryotic RNA polymerase. In a preferred embodiment, the promoter is a "strong" or "efficient" promoter. An example of a strong promoter is the immediate early human cytomegalovirus promoter (Chapman et al, 1991 *Nucl. Acids Res*19:3979-3986, which is incorporated by reference), preferably without intronic sequences. Most preferred

for use within the instant adenoviral vector is a human CMV promoter without intronic sequences, like intron A. Applicants have found that intron A, a portion of the human cytomegalovirus promoter (hCMV), constitutes a region of instability for adenoviral vectors. CMV without intron A has been found to effectuate (Examples 1-3) comparable expression capabilities in vitro when driving HIV gag expression and, furthermore, behaved equivalently to intron A-containing constructs in Balb/c mice in vivo with respect to their antibody and T-cell responses at both dosages of plasmid DNA tested (20 µg and 200 µg). Those skilled in the art will appreciate that any of a number of other known promoters, such as the strong immunoglobulin, or other eukaryotic gene promoters may also be used, including the EF1 alpha promoter, the murine CMV promoter, Rous sarcoma virus (RSV) promoter, SV40 early/late promoters and the beta-actin promoter.

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In preferred embodiments, the promoter may also comprise a regulatable sequence such as the Tet operator sequence. This would be extremely useful, for example, in cases where the gene products are effecting a result other than that desired and repression is sought.

The combination of the CMV promoter (devoid of the intron A region) with the BGH terminator is particularly preferred although other promoter/terminator combinations in the context of FG adenovirus may also be used.

Other embodiments incorporate a leader or signal peptide into the transgene. A preferred leader is that from the tissue-specific plasminogen activator protein, tPA. Examples include but are not limited to the various tPA-gag, tPA-pol and tPA-nef adenovirus-based vaccines disclosed throughout this specification.

In view of the improved adenovirus vectors described herein, an essential portion of the present invention are adenoviral-based HIV vaccines comprising said adenovirus backbones which may be administered to a mammalian host, preferably a human host, in either a prophylactic or therapeutic setting. The HIV vaccines of the present invention, whether administered alone or in combination regimens with other viral- or non-viral-based DNA vaccines, should elicit potent and broad cellular immune responses against HIV that will either lessen the likelihood of persistent virus infection and/or lead to the establishment of a clinically significant lowered virus load

subject to HIV infection or in combination with HAART therapy, mitigate the effects of previously established HIV infection (antiviral immunotherapy(ARI)). While any HIV antigen (e.g., gag, pol, nef, gp160, gp41, gp120, tat, rev, etc.) may be utilized in the herein described recombinant adenoviral vectors, preferred embodiments include the codon optimized p55 gag antigen (herein exemplified as MRKAd5gag), pol and nef. Sequences based on different Clades of HIV-1 are suitable for use in the instant invention, most preferred of which are Clade B and Clade C. Particularly preferred embodiments are those sequences (especially, codon-optimized sequences) based on concensus Clade B sequences. Preferred versions of the MRKAd5pol and MRKAd5nef series of adenoviral vaccines will encode modified versions of pol or nef, as discussed herein. Preferred embodiments of the MRKAd5HIV-1 vectors carrying HIV envelope genes and modifications thereof comprise the HIV codon-optimized env sequences of PCT International Applications PCT/US97/02294 and PCT/US97/10517, published August 28, 1997 (WO 97/31115) and December 24, 1997, respectively; both documents of which are hereby incorporated by reference.

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A most preferred aspect of the instant invention is the disclosed use of the adenoviral vector described above to effectuate expression of HIV gag. Sequences for many genes of many HIV strains are publicly available in GENBANK and primary, field isolates of HIV are available from the National Institute of Allergy and Infectious Diseases (NIAID) which has contracted with Quality Biological (Gaithersburg, MD) to make these strains available. Strains are also available from the World Health Organization (WHO), Geneva Switzerland. It is preferred that the gag gene be from an HIV-1 strain (CAM-1; Myers et al, eds. "Human Retroviruses and AIDS: 1995, IIA3-IIA19, which is hereby incorporated by reference). This gene closely resembles the consensus amino acid sequence for the clade B (North American/European) sequence. Therefore, it is within the purview of the skilled artisan to choose an appropriate nucleotide sequence which encodes a specific HIV gag antigen, or immunologically relevant portion thereof. As shown in Example 25, a clade B or clade C based p55 gag antigen will potentially be useful on a global scale. As noted herein, the transgene of choice for insertion in to a DNA or MRKAd-based adenoviral vector of the present invention is a codon optimized version of p55 gag. Such a MRKAd5gag adenoviral vector is documented in Example 11 and is at least referred to herein as MRKAd5HIV-1gag. Of course, additional versions are contemplated, including but not limited to modifications such as promoter (e.g., mCMV for hCMV) and/or pA-terminations signal (SPA for bGH) switching, as well as generating MRK Ad5 backbones with or without deletion of the Ad5 E3 gene.

The present invention also relates a series of MRKAd5pol-based adenoviral vaccines which are shown herein to generate cellular immune responses subsequent to administration in mice and non-human primate studies. Several of the MRKAd5pol series are exemplified herein. One such adenoviral vector is referred to as MRKAd5hCMV-inact opt pol(E3+), which comprises the MRKAd5 backbone, the 5 hCMV promoter (no intron A), an inactivated pol transgene, and contains the Ad5 E3 gene in the adenoviral backbone. A second exemplified pre-adenovirus plasmid and concomitant virus is referred to as MRKAd5hCMV-inact opt pol(E3-), which is identical to the former adenoviral vector except that the E3 is deleted. Both constructions contain a codon optimized, inactivated version of HIV-1 Pol, wherein at 10 least the entire coding region is disclosed herein as SEQ ID NO:3 and the expressed protein is shown as SEQ ID NO:4 (see also Figure 17A-C and Table 1, which show targeted deletion for inactivated pol. This and other preferred codon optimized versions of HIV Pol as disclosed herein are essentially as described in U.S. Application Serial No. 09/745,221, filed December 21, 2000 and PCT International 15 Application PCT/US00/34724, also filed December 21, 2000, both documents which are hereby incorporated by reference. As disclosed in the above-mentioned documents, the open reading frame for these codon-optimized HIV-1 Pol-based DNA vaccines are represented by codon optimized DNA molecules encoding codon optimized HIV-1 Pol (e.g. SEQ ID NO:2), codon optimized HIV-1 Pol fused to an 20 amino terminal localized leader sequence (e.g. SEQ ID NO:6), and especially preferable, and exemplified by the MRKAd5-Pol construct in e.g., Example 19, biologically inactivated pol ("inact opt Pol"; e.g., SEQ ID NO:4) which is devoid of significant PR, RT, RNase or IN activity associated with wild type Pol. In addition, a construct related to SEO ID NO:4 is contemplated which contains a leader peptide at 25 the amino terminal region of the IA Pol protein. A specific construct is ligated within an appropriate DNA plasmid vector containing regulatory regions operatively linked to the respective HIV-1 Pol coding region, with or without a nucleotide sequence encoding a functional leader peptide. To this end, various HIV-1 Pol constructs 30 disclosed herein relate to open reading frames for cloning to the enhanced first generation Ad vectors of the present invention (such a series of MRKAd5pol adenoviral vaccine vectors), including but not limited to wild type Pol (comprising the DNA molecule encoding WT opt Pol, as set forth in SEQ ID NO:2), tPA-opt WTPol, (comprising the DNA molecule encoding tPA Pol, as set forth in SEQ ID NO:6), inact opt Pol (comprising the DNA molecule encoding IA Pol, as set forth in SEQ ID 35 NO:4), and tPA-inact opt Pol, (comprising the DNA molecule encoding tPA-inact opt

Pol, as set forth in SEQ ID NO:8). The pol-based versions of enhanced first generation adenovirus vaccines elicit CTL and Th cellular immune responses upon administration to the host, including primates and especially humans. As noted in the above, an effect of the cellular immune-directed vaccines of the present invention should be a lower transmission rate to previously uninfected individuals and/or reduction in the levels of the viral loads within an infected individual, so as to prolong the asymptomatic phase of HIV-1 infection.

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The present invention further relates to a series of MRKAd5nef-based adenoviral vaccines which, similar to HIV gag and pol antigens, generate cellular immune responses subsequent to administration in mice and non-human primate 10 studies. The MRKAd5nef series are exemplified herein by utilizing the improved MRK adenoviral backbone in combination with modified versions of HIV nef. These exemplified MRKAd5nef vectors are as follows: (1) MRKAd5hCMVnef(G2A,LLAA) (E3+), which comprises the improved MRKAd5 backbone, a human CMV promoter an intact Ad5 E3 gene and a modified nef gene: (2) MRKAd5mCMV-15 nef(G2A,LLAA) (E3+), which is the same as (1) above but substituting a murine CMV promoter for a human CMV promoter; and (3) MRKAd5mCMV-tpanef(LLAA) (E3+), which is the same as (2) except that the nef transgene is tpanef(LLAA). Codon optimized versions of HIV-1 Nef and HIV-1 Nef modifications are essentially as described in U.S. Application Serial No. 09/738,782, filed December 15, 2000 and 20 PCT International Application PCT/US00/34162, also filed December 15, 2000, both documents which are hereby incorporated by reference. Particular embodiments of codon optimized Nef and Nef modifications relate to a DNA molecule encoding HIV-1 Nef from the HIV-1 jfrl isolate wherein the codons are optimized for expression in a mammalian system such as a human. The DNA molecule which encodes this protein 25 is disclosed herein as SEQ ID NO:9, while the expressed open reading frame is disclosed herein as SEQ ID NO:10. Another embodiment of Nef-based coding regions for use in the adenoviral vectors of the present invention comprise a codon optimized DNA molecule encoding a protein containing the human plasminogen activator (tpa) leader peptide fused with the NH2-terminus of the HIV-1 Nef 30 polypeptide. The DNA molecule which encodes this protein is disclosed herein as SEO ID NO:11, while the expressed open reading frame is disclosed herein as SEQ ID NO:12. Another modified Nef optimized coding region relates to a DNA molecule encoding optimized HIV-1 Nef wherein the open reading frame codes for modifications at the amino terminal myristylation site (Gly-2 to Ala-2) and 35 substitution of the Leu-174-Leu-175 dileucine motif to Ala-174-Ala-175, herein

described as opt nef (G2A, LLAA). The DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:13, while the expressed open reading frame is disclosed herein as SEQ ID NO:14. MRKAd5nef vectors (1) MRKAd5hCMV-nef(G2A,LLAA) (E3+) and (2) MRKAd5mCMV-nef(G2A,LLAA) (E3+) contain this transgene. An additional embodiment relates to a DNA molecule encoding optimized HIV-1 Nef wherein the amino terminal myristylation site and dileucine motif have been deleted, as well as comprising a tPA leader peptide. This DNA molecule, opt tpanef (LLAA), comprises an open reading frame which encodes a Nef protein containing a tPA leader sequence fused to amino acid residue 6-216 of HIV-1 Nef (jfrl), wherein Leu-174 and Leu-175 are substituted with Ala-174 and Ala-175, herein referred to as opt tpanef (LLAA) is disclosed herein as SEQ ID NO:15, while the expressed open reading frame is disclosed herein as SEQ ID NO:16. The MRKAd5nef vector "MRKAd5mCMV-tpanef(LLAA) (E3+)" contains this transgene.

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Along with the improved MRKAd5gag adenovirus vaccine vector described herein, generation of a MRKAd5pol and MRKAd5nef adenovirus vector provide for enhanced HIV vaccine capabilities. Namely, the generation of this trio of adenoviral vaccine vectors, all shown to generate effective cellular immune responses subsequent to host administration, provide for the ability to administer these vaccine candidates not only alone, but preferably as part of a divalent (i.e., gag and nef, gag and pol, or pol and nef components) or a trivalent vaccine (i.e., gag, pol and nef components). Therefore, a preferred aspect of the present invention are vaccine formulations and associated methods of administration and concomitant generation of host cellular immune responses associated with formulating three separate series of MRKAd5based adenoviral vector vaccines. Of course, this MRKAd5 vaccine series based on distinct HIV antigens promotes expanded opportunities for formulation of a divalent or trivalent vaccine, or possibly administration of separate formulations of one or more monovalent or divalent formulations within a reasonable window of time. It is also within the scope of the present invention to embark on combined modality regimes which include multiple but distinct components from a specific antigen. An example, but certainly not a limitation, would be separate MRKAd5pol vectors, with one vaccine vector expressing wild type Pol (SEQ ID NO:2) and another MRKAd5pol vector expressing inactivated Pol (SEQ ID NO:6). Another example might be separate MRKAd5nef vectors, with one vaccine vector expressing the tPA/LLAA version of Nef (SEQ ID NO:16) and another MRKAd5nef vector expressing the G2A,LLAA modified version of Nef (SEQ ID NO:14). Therefore, the MRKAd5 adenoviral vectors of the present invention may be used in combination

with multiple, distinct HIV antigen classes. Each HIV antigen class is subject to sequence manipulation, thus providing for a multitude of potential vaccine combinations; and such combinations are within the scope of the present invention. The utilization of such combined modalities vaccine formulation and administration increase the probability of eliciting an even more potent cellular immune response when compared to inoculation with a single modality regimen.

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The present invention also relates to application of a mono-, dual-, or trimodality administration regime of the MRKAd5gag, pol and nef adenoviral vaccine series in a prime/boost vaccination schedule. This prime/boost schedule may include any reasonable combination of the MRKAd5gag, pol and nef adenoviral vaccine series disclosed herein. In addition, a prime/boost regime may also involve other viral and/or non-viral DNA vaccines. A preferable addition to an adenoviral vaccine vector regime includes but is not limited to plasmid DNA vaccines, especially DNA plasmid vaccines that contain at least one of the codon optimized gag, pol and nef constructions, as disclosed herein.

Therefore, one aspect of this invention is the administration of the adenoviral vector containing the optimized gag gene in a prime/boost regiment in conjunction with a plasmid DNA encoding gag. To distinguish this plasmid from the adenoviralcontaining shuttle plasmids used in the construction of an adenovirus vector, this plasmid will be referred to as a "vaccine plasmid" or "DNA plasmid vaccine". Preferred vaccine plasmids for use in this administration protocol are disclosed in pending U.S. patent application 09/017,981, filed February 3, 1998 and WO98/34640, published August 13, 1998, both of which are hereby incorporated by reference. Briefly, the preferred vaccine plasmid is designated V1Jns-FLgag, which expresses the same codon-optimized gag gene as the adenoviral vectors of this invention (see Figure 2 for the nucleotide sequence of the exemplified optimized codon version of full length p55 gag). The vaccine plasmid backbone, designated V1Ins contains the CMV immediate-early (IE) promoter and intron A, a bovine growth hormone-derived polyadenylation and transcription termination sequence as the gene expression regulatory elements, and a minimal pUC backbone; see Montgomery et al., 1993, DNA Cell Biol. 12:777-783. The pUC sequence permits high levels of plasmid production in E. coli and has a neomycin resistance gene in place of an ampicillin resistance gene to provide selected growth in the presence of kanamycin. Alternatively, a vaccine plasmid which has the CMV promoter deleted of intron A can be used. Those of skill in the art will recognize that alternative vaccine plasmid

vectors may be easily substituted for these specific constructs, and this invention specifically envisions use of such alternative plasmid DNA vaccine vectors.

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Another aspect of the present invention is a prime/boost regimen which includes a vaccine plasmid which encodes an HIV pol antigen, preferably a codon optimized form of pol and also preferably a vaccine plasmid which comprises a nucleotide sequence which encodes a Pol antigen selected from the group of Pol antigens as shown in SEQ ID NOs: 2, 4, 6 and 8. The variety of potential DNA plasmid vaccines which encode various biologically active forms of HIV-1 Pol, wherein administration, intracellular delivery and expression of the HIV-1 Pol gene of interest elicits a host CTL and Th response. The preferred synthetic DNA molecules of the present invention encode codon optimized wild type Pol (without Pro activity) and various codon optimized inactivated HIV-1 Pol proteins. The HIV-1 pol open reading disclosed herein are especially preferred for pharmaceutical uses, especially for human administration as delivered via a recombinant adenoviral vaccine, especially an enhanced first generation recombinant adenoviral vaccine as described herein. Several embodiments of this portion of the invention are provided in detail below, namely DNA molecules which comprise a HIV-1 pol open reading frame, whether encoding full length pol or a modification or fusion as described herein, wherein the codon usage has been optimized for expression in a mammal, especially a human. Again, these DNA sequences are positioned appropriately within a recombinant adenoviral vector, such as the exemplified recombinant adenoviral vector described herein, so as to promote expression of the respective HIV-1 Pol gene of interest, and subsequent to administration, elicit a host CTL and Th response. Again, these preferred, but in no way limiting, pol genes are as disclosed herein and essentially as described in U.S. Application Serial No. 09/745,221, filed December 21, 2000 and PCT International Application PCT/US00/34724, also filed December 21, 2000, both documents which are hereby incorporated by reference.

A third series of vaccine plasmids which are useful in a combined modality and/or prime/boost regimen are vaccine plasmids which encode an HIV nef antigen or biologically and/or immunologically relevant modification thereof. As noted elsewhere, preferred vaccine plasmids contain a codon optimized form of nef and also preferably comprise a nucleotide sequence which encodes a Nef antigen selected from the group of Nef antigens as shown in SEQ ID NOs: 10, 12, 14 and 16. These preferred nef coding regions are disclosed herein, as well as being described in U.S. Application Serial No. 09/738,782, filed December 15, 2000 and PCT International

Application PCT/US00/34162, also filed December 15, 2000, both documents which are hereby incorporated by reference.

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Therefore, the adenoviral vaccines and plasmid DNA vaccines of this invention may be administered alone, or may be part of a prime and boost administration regimen. A mixed modality priming and booster inoculation scheme will result in an enhanced immune response, particularly is pre-existing anti-vector immune responses are present. This one aspect of this invention is a method of priming a subject with the plasmid vaccine by administering the plasmid vaccine at least one time, allowing a predetermined length of time to pass, and then boosting by administering the adenoviral vaccine. Multiple primings typically, 1-4, are usually employed, although more may be used. The length of time between priming and boost may typically vary from about four months to a year, but other time frames may be used. In experiments with rhesus monkeys, the animals were primed four times with plasmid vaccines, then were boosted 4 months later with the adenoviral vaccine. Their cellular immune response was notably higher than that of animals which had only received adenoviral vaccine. The use of a priming regimen may be particularly preferred in situations where a person has a pre-existing anti-adenovirus immune response.

Furthermore and in the alternative, multiple HIV-1 viral antigens, such as the MRKAd5 adenoviral vaccines disclosed herein, may be ligated into a proper shuttle plasmid for generation of a pre-adenoviral plasmid comprising multiple open reading frames. For example a trivalent vector may comprise a gag-pol-nef fusion, in either a E3(-) or E3(+) background, preferably a E3 deleted backbone, or possible a "2+1" divalent vaccine, such as a gag-pol fusion (i.e., codon optimized p55 gag and inactivated optimized pol; Example 29 and Table 25) within the same MRKAd5 backbone, with each open reading frame being operatively linked to a distinct promoter and transcription termination sequence. Alternatively, the two open reading frames may be operatively linked to a single promoter, with the open reading frames operatively linked by an internal ribosome entry sequence (IRES), as disclosed in International Publication No. WO 95/24485, which is hereby incorporated by reference. Figure 9 shows that the use of multiple promoters and termination sequences provide for similar growth properties, while Figure 28 shows that these MRKAd5gag-based vectors are also stable at least through passage 21. In the absence of the use of IRES-based technology, it is preferred that a distinct promoter be used to support each respective open reading frame, so as to best preserve vector stability. As examples, and certainly not as limitations, potential multiple transgene vaccines may

include a three transgene vector such as hCMV-gagpol-bGHpA + mCMV-nef-SPA in an E3 deleted backbone or hCMV-gagpol-bGHpA + mCMV-nef-SPA(E3+). Potential "2+1" divalent vaccines of the present invention might be a hCMV-gagbGHpA + mCMV-nef-SPA in an E3+ backbone (vector #1) in combination with hCMV-pol-bGHpA in an E3+ backbone (vector #2), with all transgenes in the E1 parallel orientation. Fusion constructs other than the gag-pol fusion described above are also suitable for use in various divalent vaccine strategies and can be composed of any two HIV antigens fused to one another (e.g.,, nef-pol and gag-nef). These adenoviral compositions are, as above, preferably delivered along with an adenoviral composition comprising an additional HIV antigen in order to diversify the immune response generated upon administration. Therefore, a multivalent vaccine delivered in a single, or possible second, adenoviral vector is certainly contemplated as part of the present invention. Again, this mode of administration is another example of whereby an efficaceous adenovirus-based HIV-1 vaccine may be administered via a combined modality regime. It is important to note, however, that in terms of deciding on an insert for the disclosed adenoviral vectors, due consideration must be dedicated to the effective packaging limitations of the adenovirus vehicle. Adenovirus has been shown to exhibit an upper cloning capacity limit of approximately 105% of the wildtype Ad5 sequence.

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Regardless of the gene chosen for expression, it is preferred that the sequence be "optimized" for expression in a human cellular environment. A "triplet" codon of four possible nucleotide bases can exist in 64 variant forms. That these forms provide the message for only 20 different amino acids (as well as transcription initiation and termination) means that some amino acids can be coded for by more than one codon. Indeed, some amino acids have as many as six "redundant", alternative codons while some others have a single, required codon. For reasons not completely understood, alternative codons are not at all uniformly present in the endogenous DNA of differing types of cells and there appears to exist variable natural hierarchy or "preference" for certain codons in certain types of cells. As one example, the amino acid leucine is specified by any of six DNA codons including CTA, CTC, CTG, CTT, TTA, and TTG (which correspond, respectively, to the mRNA codons, CUA, CUC, CUG, CUU, UUA and UUG). Exhaustive analysis of genome codon frequencies for microorganisms has revealed endogenous DNA of E. coli most commonly contains the CTG leucine-specifying codon, while the DNA of yeasts and slime molds most commonly includes a TTA leucine-specifying codon. In view of this hierarchy, it is generally held that the likelihood of obtaining high levels of expression of a leucine-

rich polypeptide by an *E. coli* host will depend to some extent on the frequency of codon use. For example, a gene rich in TTA codons will in all probability be poorly expressed in *E. coli*, whereas a CTG rich gene will probably highly express the polypeptide. Similarly, when yeast cells are the projected transformation host cells for expression of a leucine-rich polypeptide, a preferred codon for use in an inserted DNA would be TTA.

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The implications of codon preference phenomena on recombinant DNA techniques are manifest, and the phenomenon may serve to explain many prior failures to achieve high expression levels of exogenous genes in successfully transformed host organisms—a less "preferred" codon may be repeatedly present in the inserted gene and the host cell machinery for expression may not operate as efficiently. This phenomenon suggests that synthetic genes which have been designed to include a projected host cell's preferred codons provide a preferred form of foreign genetic material for practice of recombinant DNA techniques. Thus, one aspect of this invention is an adenovirus vector or adenovirus vector in some combination with a vaccine plasmid where both specifically include a gene which is codon optimized for expression in a human cellular environment. As noted herein, a preferred gene for use in the instant invention is a codon-optimized HIV gene and, particularly, HIV gag, pol or nef.

Adenoviral vectors in accordance with the instant invention can be constructed using known techniques, such as those reviewed in Hitt et al, 1997 "Human Adenovirus Vectors for Gene Transfer into Mammalian Cells" Advances in Pharmacology 40:137-206, which is hereby incorporated by reference.

In constructing the adenoviral vectors of this invention, it is often convenient to insert them into a plasmid or shuttle vector. These techniques are known and described in Hitt et al., *supra*. This invention specifically includes both the adenovirus and the adenovirus when inserted into a shuttle plasmid.

Preferred shuttle vectors contain an adenoviral portion and a plasmid portion. The adenoviral portion is essentially the same as the adenovirus vector discussed supra, containing adenoviral sequences (with non-functional or deleted E1 and E3 regions) and the gene expression cassette, flanked by convenient restriction sites. The plasmid portion of the shuttle vector often contains an antibiotic resistance marker under transcriptional control of a prokaryotic promoter so that expression of the antibiotic does not occur in eukaryotic cells. Ampicillin resistance genes, neomycin resistance genes and other pharmaceutically acceptable antibiotic resistance markers may be used. To aid in the high level production of the polynucleotide by

fermentation in prokaryotic organisms, it is advantageous for the shuttle vector to contain a prokaryotic origin of replication and be of high copy number. A number of commercially available prokaryotic cloning vectors provide these benefits. It is desirable to remove non-essential DNA sequences. It is also desirable that the vectors not be able to replicate in eukaryotic cells. This minimizes the risk of integration of polynucleotide vaccine sequences into the recipients' genome. Tissue-specific promoters or enhancers may be used whenever it is desirable to limit expression of the polynucleotide to a particular tissue type.

In one embodiment of this invention, the pre-plasmids (e.g., pMRKAd5pol, pMRKAd5nef and pMRKAd5gag were generated by homologous recombination using the MRKHVE3 (and MRKHVO for the E3- version) backbones and the appropriate shuttle vector, as shown for pMRKAd5pol in Figure 22 and for pMRKAd5nef in Figure 23. The plasmid in linear form is capable of replication after entering the PER.C6[®] cells and virus is produced. The infected cells and media were harvested after viral replication was complete.

Viral vectors can be propagated in various E1 complementing cell lines, including the known cell lines 293 and PER.C6[®]. Both these cell lines express the adenoviral E1 gene product. PER.C6[®] is described in WO 97/00326 (published January 3, 1997) and issued U.S. Patent No. 6,033,908, both of which are hereby incorporated by reference. It is a primary human retinoblast cell line transduced with an E1 gene segment that complements the production of replication deficient (FG) adenovirus, but is designed to prevent generation of replication competent adenovirus by homologous recombination. Cells of particular interest have been stably transformed with a transgene that encodes the AD5E1A and E1B gene, like PER.C6[®], from 459 bp to 3510 bp inclusive. 293 cells are described in Graham et al., 1977 J. Gen. Virol 36:59-72, which is hereby incorporated by reference. As stated above, consideration must be given to the adenoviral sequences present in the complementing cell line used. It is important that the sequences not overlap with that present in the vector if the possibility of recombination is to be minimized.

It has been found that vectors generated in accordance with the above description are more effective in inducing an immune response and, thus, constitute very promising vaccine candidates. More particularly, it has been found that first generation adenoviral vectors in accordance with the above description carrying a codon-optimized HIV gag gene, regulated with a strong heterologous promoter can be used as human anti-HIV vaccines, and are capable of inducing immune responses.

Standard techniques of molecular biology for preparing and purifying DNA constructs enable the preparation of the DNA immunogens of this invention.

A vaccine composition comprising an adenoviral vector in accordance with the instant invention may contain physiologically acceptable components, such as buffer, normal saline or phosphate buffered saline, sucrose, other salts and polysorbate. One preferred formulation has: 2.5-10 mM TRIS buffer, preferably about 5 mM TRIS buffer; 25-100 mM NaCl, preferably about 75 mM NaCl; 2.5-10% sucrose, preferably about 5% sucrose; 0.01 -2 mM MgCl₂; and 0.001%-0.01% polysorbate 80 (plant derived). The pH should range from about 7.0-9.0, preferably about 8.0. One skilled in the art will appreciate that other conventional vaccine excipients may also be used it make the formulation. The preferred formulation contains 5mM TRIS, 75 mM NaCl, 5% sucrose, 1mM MgCl₂, 0.005% polysorbate 80 at pH 8.0 This has a pH and divalent cation composition which is near the optimum for Ad5 stability and minimizes the potential for adsorption of virus to a glass surface. It does not cause tissue irritation upon intramuscular injection. It is preferably frozen until use.

The amount of adenoviral particles in the vaccine composition to be introduced into a vaccine recipient will depend on the strength of the transcriptional and translational promoters used and on the immunogenicity of the expressed gene product. In general, an immunologically or prophylactically effective dose of $1x10^7$ to $1x10^{12}$ particles and preferably about $1x10^{10}$ to $1x10^{11}$ particles is administered directly into muscle tissue. Subcutaneous injection, intradermal introduction, impression through the skin, and other modes of administration such as intraperitoneal, intravenous, or inhalation delivery are also contemplated. It is also contemplated that booster vaccinations are to be provided. Following vaccination with HIV adenoviral vector, boosting with a subsequent HIV adenoviral vector and/or plasmid may be desirable. Parenteral administration, such as intravenous, intramuscular, subcutaneous or other means of administration of interleukin-12 protein, concurrently with or subsequent to parenteral introduction of the vaccine compositions of this invention is also advantageous.

The adenoviral vector and/or vaccine plasmids of this invention polynucleotide may be unassociated with any proteins, adjuvants or other agents which impact on the recipients' immune system. In this case, it is desirable for the vector to be in a physiologically acceptable solution, such as, but not limited to, sterile saline or sterile buffered saline. Alternatively, the vector may be associated with an adjuvant known in the art to boost immune responses (i.e., a "biologically effective"

adjuvant), such as a protein or other carrier. Vaccine plasmids of this invention may, for instance, be delivered in saline (e.g., PBS) with or without an adjuvant. Preferred adjuvants are Alum or CRL1005 Block Copolymer. Agents which assist in the cellular uptake of DNA, such as, but not limited to, calcium ions, may also be used to advantage. These agents are generally referred to herein as transfection facilitating reagents and pharmaceutically acceptable carriers. Techniques for coating microprojectiles coated with polynucleotide are known in the art and are also useful in connection with this invention.

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This invention also includes a prime and boost regimen wherein a first adenoviral vector is administered, then a booster dose is given. The booster dose may be repeated at selected time intervals. Alternatively, a preferred inoculation scheme comprises priming with a first adenovirus serotype and then boosting with a second adenovirus serotype. More preferably, the inoculation scheme comprises priming with a first adenovirus serotype and then boosting with a second adenovirus serotype, wherein the first and second adenovirus serotypes are classified within separate subgroups of adenoviruses. The above prime/boost schemes are particularly preferred in those situations where a preexisting immunity is identified to the adenoviral vector of choice. In this type of scheme, the individual or population of individuals is primed with an adenovirus of a serotype other than that to which the preexisting immunity is identified. This enables the first adenovirus to effectuate sufficient expression of the transgene while evading existing immunity to the second adenovirus (the boosting adenovirus) and, further, allows for the subsequent delivery of the transgene via the boosting adenovirus to be more effective. Adenovirus serotype 5 is one example of a virus to which such a scheme might be desirable. In accordance with this invention, therefore, one might decide to prime with a non-group C adenovirus (e.g., Ad12, a group A adenovirus, Ad24, a group D adenovirus, or Ad35, a group B adenovirus) to evade anti-Ad5 immunity and then boost with Ad5, a group C adenovirus. Another preferred embodiment involves administration of a different adenovirus (including non-human adenovirus) vaccine followed by administration of the adenoviral vaccines disclosed. In the alternative, a viral antigen of interest can be first delivered via a viral vaccine other than an adenovirus-based vaccine, and then followed with the adenoviral vaccine disclosed. Alternative viral vaccines include but are not limited to pox virus and venezuelan equine encephilitis virus.

A large body of human and animal data supports the importance of cellular immune responses, especially CTL in controlling (or eliminating) HIV infection. In humans, very high levels of CTL develop following primary infection and correlate

with the control of viremia. Several small groups of individuals have been described who are repeatedly exposed to HIV by remain uninfected; CTL has been noted in several of these cohorts. In the SIV model of HIV infection, CTL similarly develops following primary infection, and it has been demonstrated that addition of anti-CD8 monoclonal antibody abrogated this control of infection and leads to disease progression. This invention uses adenoviral vaccines alone or in combination with plasmid vaccines to induce CTL.

The following non-limiting Examples are presented to better illustrate the invention.

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EXAMPLE 1

Removal of the Intron A Portion of the hCMV Promoter GMP grade pVIJnsHIVgag was used as the starting material to amplify the hCMV promoter. PVIInsHIVgag is a plasmid comprising the CMV immediate-early (IE) promoter and intron A, a full-length codon-optimized HIV gag gene, a bovine growth hormone-derived polyadenylation and transcriptional termination sequence, and a minimal pUC backbone; see Montgomery et al., supra for a description of the plasmid backbone. The amplification was performed with primers suitably positioned to flank the hCMV promoter. A 5' primer was placed upstream of the Msc1 site of the hCMV promoter and a 3' primer (designed to contain the BgIII recognition sequence) was placed 3' of the hCMV promoter. The resulting PCR product (using high fidelity Taq polymerase) which encompassed the entire hCMV promoter (minus intron A) was cloned into TOPO PCR blunt vector and then removed by double digestion with Msc1 and BgIII. This fragment was then cloned back into the original GMP grade pV1JnsHIVgag plasmid from which the original promoter, intron A, and the gag gene were removed following Msc1 and BgIII digestion. This ligation reaction resulted in the construction of a hCMV promoter (minus intron A) + bGHpA expression cassette within the original pV1JnsHIVgag vector backbone. This vector is designated pVIInsCMV(no intron).

The FLgag gene was excised from pV1JnsHIVgag using BgIII digestion and the 1,526 bp gene was gel purified and cloned into pV1JnsCMV(no intron) at the BgIII site. Colonies were screened using Sma1 restriction enzymes to identify clones that carried the Flgag gene in the correct orientation. This plasmid, designated pV1JnsCMV(no intron)-FLgag-bGHpA, was fully sequenced to confirm sequence integrity.

The plasmid, pV1Jns-mCMV-FLgag-bGHpA, is identical to the pV1JnsCMV(no intron)-FLgag-bGHpA except that the hCMV promoter has been removed and replaced with the murine CMV (mCMV) promoter.

Figure 3 diagrammatically shows the new transgene constructs in comparison with the original transgene.

15 EXAMPLE 2

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Gag Expression Assay for Modified Gag Transgenes
Gag Elisa was performed on culture supernatants obtained from transient tissue culture transfection experiments in which the two new hCMV-containing plasmid constructs, pV1JnsCMV(no intron)-FLgag-bGHpA and pV1JnsCMV(no intron)-FLgag-SPA, both devoid of intron A, were compared to pV1JnsHIVgag which, as noted above possesses the intron A as part of the hCMV promoter. Table 2 below shows the *in vitro* gag expression data of the new gag plasmids compared with the GMP grade original plasmid. The results displayed in Table 2 show that both of the new hCMV gag plasmid constructs have expression capacities comparable to the original plasmid construct which contains the intron A portion of the hCMV promoter.

Table 2: In vitro DNA transfection of original and new plasmid HIV-1 gag constructs.

Plasmid	μg gag/10e6 COS cells/5μg DNA/48 hr
HIVFL-gagPR9901 ^a	10.8
PVIIns-hCMV-FLgag-bGHpAb	16.6
pV1Jns-hCMV-FLgag-SPA ^{b,c}	12.0

^a GMP grade pV1Ins-hCMVintronA-FLgag-bGHpA.

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EXAMPLE 3

Rodent (Balb/c) Study for Modified gag Transgenes
A rodent study was performed on the two new plasmid constructs
described above – pV1JnsCMV(no intron)-FLgag-bGHpA and pV1JnsCMV(no
intron)-FLgag-SPA - in order to compare them with the construct described above
possessing the intron A portion of the CMV promoter, pV1JnsHIVgag. Gag antibody
and Elispot responses (described in PCT International Application No.
PCT/US00/18332 (WO 01/02607) filed July 3, 2000, claiming priority to U.S.
Provisional Application Serial No. 60/142,631, filed July 6, 1999 and U.S.
Application Serial No. 60/148,981, filed August 13, 1999, all three applications which
are hereby incorporated by reference) were measured. The results displayed in Table
3 below, show that the new plasmid constructs behaved equivalently to the original
construct in Balb/c mice with respect to their antibody and T-cell responses at both
dosages of plasmid DNA tested, 20 µg and 200 µg.

⁵ b New plasmid constructions that have the intron A portion removed from the hCMV promoter.

^c In this construct the bGH terminator has been replaced with the short synthetic polyadenylation signal (SPA)

EXAMPLE 4

Table 3: HIV191: Immunogenicity of V1Jns-gag under different promoter and termination control elements.

DNA*	Dose,	Anti-p24 Titers			SFC/10^6 Cells		
	ug ^b	(3 Wk PD1) ^c			(4 Wk PD1) ^d		
Promoter/terminator	-	GMT	+SE_	-SE	Media	gag197-205	p24
HIVFL-gagPR9901	200	12800	4652	3412	2(2)	129(19)	30(11)
(GMP grade)	20	5572	1574	1227	0	56(9)	25(6)
pV1Jns-hCMV-	200	11143	2831	2257	0	98(5)	12(6)
FL-gag-bGHpA	20	7352	2808	2032		73(9)	11(6)
pV1Jns-hCMV-	200	16890	5815	4326	1(1)	94(4)	26(7)
FL-gag-SPA	20	5971	5361	2825	0	85(17)	38(10)
Naīve	0	123	. 50	36	0	0	0

in PBS

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Construction of the Modified Shuttle Vector - "MRKpdelE1 Shuttle"

The modifications to the original Ad5 shuttle vector (pdelE1sp1A; a vector comprising Ad5 sequences from basepairs 1-341 and 3524-5798, with a multiple cloning region between nucleotides 341 and 3524 of Ad5, included the following three manipulations carried out in sequential cloning steps as follows:

- (1) The left ITR region was extended to include the *Pac1* site at the junction between the vector backbone and the adenovirus left ITR sequences. This allow for easier manipulations using the bacterial homologous recombination system.
- 10 (2) The packaging region was extended to include sequences of the wild-type (WT) adenovirus from 342 bp to 450 bp inclusive.
 - (3) The area downstream of pIX was extended 13 nucleotides (i.e., nucleotides 3511-3523 inclusive).

These modifications (Figure 4) effectively reduced the size of the E1 deletion without overlapping with any part of the E1A/E1B gene present in the transformed PER.C6[®] cell line. All manipulations were performed by modifying the Ad shuttle vector pdelE1sp1A.

Once the modifications were made to the shuttle vector, the changes were incorporated into the original Ad5 adenovector backbones (pAdHVO and pAdHVE3) by bacterial homologous recombination using *E. coli* BJ5183 chemically competent cells.

bi.m. Injections into both quads, 50 μL per quad

cn=10;GMT, geometric mean titer, SE, standard. error

dn=5, pooled spleens; mean of triplicate wells and standard, deviation, in parentheses;

EXAMPLE 5

Construction of Modified Adenovector Backbones (E3+ and E3-)

The original adenovectors pAdHVO (comprising all Ad5 sequences except those nucleotides encompassing the E1 and E3 regions) and pADHVE3 (comprising all Ad5 sequences except those nucleotides encompassing the E1 region), were each 5 reconstructed so that they contained the modifications to the E1 region. This was accomplished by digesting the newly modified shuttle vector (MRKpdelE1 shuttle) with Pac1 and BstZ1101 and isolating the 2,734 bp fragment which corresponds to the adenovirus sequence. This fragment was co-transformed with DNA from either Cla1 linearized pAdHVO (E3- adenovector) or Cla1 linearized pAdHVE3 10 (E3+adenovector) into E. coli BJ5183 competent cells. At least two colonies from each transformation were selected and grown in Terrific™ broth for 6-8 hours until turbidity was reached. DNA was extracted from each cell pellet and then transformed into E. coli XL1 competent cells. One colony from each transformation was selected and grown for plasmid DNA purification. The plasmid was analyzed by restriction 15 digestions to identify correct clones. The modified adenovectors were designated MRKpAdHVO (E3- plasmid) and MRKpAdHVE3 (E3+ plasmid). Virus from these new adenovectors (MRKHVO and MRKHVE3, respectively) as well as the old version of the adenovectors were generated in the PER.C6® cell lines to accommodate the following series of viral competition experiments. In addition, the multiple 20 cloning site of the original shuttle vector contained ClaI, BarnHI, Xho I, EcoRV, HindIII, Sal I, and Bgl II sites. This MCS was replaced with a new MCS containing Not I, Cla I, EcoRV and Asc I sites. This new MCS has been transferred to the MRKpAdHVO and MRKpAdHVE3 pre-plasmids along with the modification made to the packaging region and pIX gene. 25

EXAMPLE 6

Analysis of the Effect of the Packaging Signal Extension

To study the effects of the modifications made to the E1 deletion region, the viruses obtained from the original backbone (pAdHVE3) and the new backbone (MRKpAdHVE3) were mixed together in equal MOI ratios (1:1 and 5:5) and passaged through several rounds; see Figure 5, Expt.#1. Both of the viruses in the experiment contained the E3 gene intact and did not contain a transgene. The only difference between the two viruses was within the region of the E1 deletion. Following the coinfection of the viruses at P1 (passage 1), the mixtures were propagated through an additional 4 passages at which time the cells were harvested

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and the virus extracted and purified by CsCl banding. The viral DNA was extracted and digested with *Hind*III and the digestion products were then radioactively labeled. For the controls, the respective pre-plasmids (pAdHVE3 ("OLD E3+"); MRKpAdHVE3 ("NEW E3+")) were also digested with *Hind*III (and *Pac1* to remove the vector backbone) and subsequently labeled with [³³P]dATP. The radioactively labeled digestion products were subjected to gel electrophoresis and the gel was dried down onto Whatman paper before being exposed to autoradiographic film. Figure 6 clearly shows that the new adenovirus which has the addition made to the packaging signal region has a growth advantage compared with the original adenovirus. In the experiments performed (at either ratio tested), only the digestion bands pertaining to the newly modified virus were present. The diagnostic band of size 3,206 (from the new virus) was clearly present. However, there was no evidence of the diagnostic band of size 2,737 bp expected from the original virus.

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EXAMPLE 7

Analysis of the Effect of the E3 Gene

The second set of the virus competition study involved mixing equal MOI ratio (1:1) of the newly modified viruses, that obtained from MRKpAdHVO and MRKpAdHVE3 (Figure 5, Expt. #2). In this set, both viruses had the new modifications made to the E1 deletion. The first virus (that from MRKpAdHVO) does not contain an E3 gene. The second virus (that from MRKpAdHVE3) does contain the E3 gene. Neither of the viruses contain a transgene. Following coinfection of the viruses, the mixtures were propagated through an additional 4 passages at which time the cells were harvested and the total virus extracted and purified by CsCl banding. The viral DNA was extracted and digested with HindIII and the digestion products were then radioactively labeled. For the controls, the respective pre-plasmids MRKpAdHVO ("NEW E3-"); MRKpAdHVE3 ("NEW E3+") were also digested with HindIII (and Pac1 to remove the vector backbone) and then labeled with [33P]dATP. The radioactively labeled digestion products were subjected to gel electrophoresis and the gel was dried down onto Whatman paper before being exposed to autoradiographic film. Figure 6 shows the results of the viral DNA analysis of the E3+ virus and E3- virus mixing experiment. The diagnostic band corresponding to the E3+ virus (5,665 bp) was present in greater amount compared with the diagnostic band of 3,010 bp corresponding to the E3- virus. This indicates that the virus that contains the E3 gene is able to amplify more rapidly

compared with the virus that does not contain an E3 gene. This increased amplification capacity has been confirmed by growth studies; see Table 4 below.

EXAMPLE 8

Construction of the new shuttle vector containing modified gag transgene – "MRKpdelE1-CMV(no intron)-FLgag-bGHpA"

The modified plasmid pV1JnsCMV(no intron)-FLgag-bGHpA was digested with Msc1 overnight and then digested with Sfi1 for 2 hours at 50°C. The DNA was then treated with Mungbean nuclease for 30 mins at 30°C. The DNA mixture was desalted using the Qiaex II kit and then Klenow treated for 30 mins at 37°C to fully blunt the ends of the transgene fragment. The 2,559 bp transgene fragment was then gel purified. The modified shuttle vector (MRKpdelE1 shuttle) was linearized by digestion with EcoRV, treated with calf intestinal phosphatase and the resulting 6,479 bp fragment was then gel purified. The two purified fragments were then ligated together and several dozen clones were screened to check for insertion of the transgene within the shuttle vector. Diagnostic restriction digestion was performed to identify those clones carrying the transgene in the E1 parallel and E1 anti-parallel orientation. This strategy was followed to clone in the other gag transgenes in the MRKpdelE1 shuttle vector.

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EXAMPLE 9

Construction of the MRK FG Adenovectors

The shuttle vector containing the HIV-1 gag transgene in the E1 parallel orientation, MRKpdelE1-CMV(no intron)-FLgag-bGHpA, was digested with Pac1. The reaction mixture was digested with BsfZ171. The 5,291 bp fragment was purified by gel extraction. The MRKpAdHVE3 plasmid was digested with Cla1 overnight at 37°C and gel purified. About 100 ng of the 5,290 bp shuttle +transgene fragment and ~100 ng of linearized MRKpAdHVE3 DNA were co-transformed into E. coli BJ5183 chemically competent cells. Several clones were selected and grown in 2 ml Terrific™ broth for 6-8 hours, until turbidity was reached. The total DNA from the cell pellet was purified using Qiagen alkaline lysis and phenol chloroform method. The DNA was precipitated with isopropanol and resuspended in 20 μl dH₂0. A 2 μl aliquot of this DNA was transformed into E. coli XL-1 competent cells. A single colony from each separate transformation was selected and grown overnight in 3 ml LB +100 μg/ml ampicillin. The DNA was isolated using Qiagen columns. A positive clone was identified by digestion with the restriction enzyme BstEII which cleaves

within the gag gene as well as the plasmid backbone. The pre-plasmid clone is designated MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA and is 37,498 bp in size. This strategy was followed to generate E3- and E3+ versions of each of the other gag transgene constructions in both E1 parallel and E1 anti-parallel versions. Figures 7A, 7B and 7C show the various combinations of adenovectors constructed.

EXAMPLE 10

Plasmid Competition Studies

A series of plasmid competition studies was carried out. Briefly, the screening of the various combinations of new constructs was performed by mixing equal amounts of each of two competing plasmids. In the experiment shown in Figure 8A, plasmids containing the same transgene but in different orientations were mixed together to create a "competition" between the two plasmids. The aim was to look at the effects of transgene orientation. In the experiment shown in Figure 8B, plasmids containing different polyadenylation signals (but in the same orientation) were mixed together in equal amounts. The aim was to assess effects of polyA signals. Following the initial transfection, the virus was passaged through ten rounds and the viral DNA analyzed by radioactive restriction analysis.

Analysis of the viral species from the plasmid mixing experiment (Figure 8A) showed that adenovectors which had the transgene inserted in the E1 parallel orientation amplified better and were able to out-compete the adenovirus which had the transgene inserted in the E1 anti-parallel orientation. Viral DNA analysis of the mixtures at passage 3 and certainly at passage 6, showed a greater ratio of the virus carrying the transgene in the E1 parallel orientation compared with the E1 antiparallel version. By passage 10, the only viral species observed was the adenovector with the transgene in the E1 parallel orientation for both transgenes tested (hCMV(no intron)-FLgag-bGHpA and hCMV(no intron)-FLgag-SPA).

Analysis of the viral species from the plasmid mixing experiment #2 (Figure 8B) at passages 3 and 6 showed that the polyadenylation signals tested (bGHpA and SPA) did not have an effect on the growth of the virus. Even at passage 10 the two viral species in the mixture were still present in equal amounts.

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EXAMPLE 11

Virus generation of an enhanced adenoviral construct - "MRK Ad5 HIV-1gag"

The results obtained from the competition study allowed us to make the following conclusions: (1) The packaging signal extension is beneficial; (2) Presence of E3 does enhance viral growth; (3) E1 parallel orientation is recommended; and (4) PolyA signals have no effect on the growth of the adenovirus.

MRK Ad5 HIV-1 gag exhibited the most desirable results. This construct contains the hCMV(no intron)-FLgag-bGHpA transgene inserted into the new E3+adenovector backbone, MRKpAdHVE3, in the E1 parallel orientation. We have designated this adenovector MRK Ad5 HIV-1 gag. This construct was prepared as outlined below:

The pre-plasmid MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA was digested was Pac1 to release the vector backbone and 3.3 µg was transfected by calcium phosphate method (Amersham Pharmacia Biotech.) in a 6 cm dish containing PER.C6[®] cells at ~60% confluence. Once CPE was reached (7-10 days), the culture was freeze/thawed three times and the cell debris pelleted. 1 ml of this cell lysate was used to infect into a 6 cm dish containing PER.C6® cells at 80-90% confluence. Once CPE was reached, the culture was freeze/thawed three times and the cell debris pelleted. The cell lysate was then used to infect a 15 cm dish containing PER.C6[®] cells at 80-90% confluence. This infection procedure was continued and expanded at passage 6. The virus was then extracted from the cell pellet by CsCl method. Two bandings were performed (3-gradient CsCl followed by a continuous CsCl gradient). Following the second banding, the virus was dialyzed in A105 buffer. Viral DNA was extracted using pronase treatment followed by phenol chloroform. The viral DNA was then digested with *Hind*III and radioactively labeled with [33P]dATP. Following gel electrophoresis to separate the digestion products the gel was dried down on Whatman paper and then subjected to autoradiography. The digestion products were compared with the digestion products from the pre-plasmid (that had been digested with Pac1/HindIII prior to labeling). The expected sizes were observed, indicating that the virus had been successfully rescued. This strategy was used to rescue virus from each of the various adenovector plasmid constructs prepared.

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EXAMPLE 12

Stability Analyses

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To determine whether the various adenovector constructs (e.g., MRK Ad5 HIV-1 gag) show genetic stability, the viruses were each passaged continually. The viral DNA was analyzed at passages 3, 6 and 10. Each virus maintained its correct genetic structure. In addition, the stability of the MRK Ad5 HIV-1 gag was analyzed under propagation conditions similar to that performed in large scale production. For this analysis, the transfections of MRK Ad5 HIV-1 gag as well as three other adenoviral vectors were repeated and the virus was purified at P3. The three other adenovectors were as follows: (1) that comprising hCMV(no intron)-Flgag with a bGHpA terminator in an E3- adenovector backbone; (2) that comprising hCMV(no intron)-Flgag with a SPA termination signal in an E3+ adenovector backbone, and that comprising a mCMV-Flgag with a bGHpA terminator in an E3+ adenovector backbone. All of the vectors have the transgene inserted in the E1 parallel orientation. Viral DNA was analyzed by radioactive restriction analysis to confirm that it was correct before being delivered to fermentation cell culture for continued passaging in serum-free media. At P5 each of the four viruses were purified and the viral DNA extracted for analysis by the restriction digestion and radiolabeling procedure. This virus has subsequently been used in a series of studies (in vitro gag expression in COS cells, rodent study and rhesus monkey study) as will be described below. The viruses 20 from P5 are shown in Figure 9.

The passaging under serum-free conditions was continued for the MRKHVE3 (transgene-less, obtained from MRKpAdHVE3 pre-plasmid) and the MRKAd5HIV-1gag (obtained from MRKpAdHVE3+CMV(no intron)-FLgagbGHpA pre-plasmid) viruses. Figure 10 shows viral DNA analysis by radioactive restriction digestion at passage 11 for MRKHVE3, MRKAd5HIV-1gagE3-, and passage 11 and 12 for MRKAd5HIV-1gag. Aside from the first lane which is the DNA marker lane, the next three lanes are virus from the pre-plasmid controls (controls based on the original virus) - MRKpAdHVE3 (also referred to as "pMRKHVE3"), MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA, and pMRKAd5gag(E3-), respectively. As seen in Figure 10, each of the viral DNA samples show the expected bands with no extraneous bands showing. This signifies that there are no major variant adenovirus species present that can be detected by autoradiography.

Figure 11 shows the results of viral competition study between MRKHVE3 and MRKAd5HIV-1gag. These viruses were mixed together at equal MOI (140 viral

particles each; 280 vp total) at passage 6 and continued to be passaged until P11. Aside from the first lane which is the DNA marker lane, the next two lanes are the pre-plasmid controls obtained from MRKpAdHVE3 and MRKpAdHVE3+CMV(no intron)-FLgag-bGHpA. The next two lanes are the viral DNA from the starting viral material at passage six. The last two lanes are the competition studies performed in duplicate. The data in Figure 11 shows the effect the gag transgene in culture. Growth of a MRKAd5gag virus was compared with growth of a "transgene-less" MRKHVE3. These two viruses were infected at the same MOI (i.e. 140 vp each) at passage 6 and then passaged through to passage 11 and the viral pool was analyzed by radioactive restriction analysis. The data shows that one virus did not out compete the other. Therefore, the gag transgene did not show obvious signs of toxicity to the adenovirus.

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Analysis by *Hind*III digestion shows that each virus specie is present in approximately equal amounts. As above, there does not appear to be signs of any extraneous bands. Figure 12 shows higher passage numbers for MRKAd5HIV-1gag grown under serum-containing conditions. The genome integrity again has been maintained and there is no evidence of rearrangements, even at the highest passage level (P21).

Each of the four vectors shown in Figure 9 were analyzed for amplification capacity. Table 4 below shows the QPA analysis used in the estimation of viral amplification ratios at P4. The determination of the amplification ratio for the original HIV-1 gag construct is based on the clinical lot at P12. It has been shown that amplification rates increases with higher passage number for the original virus. The reason for this observation is due to the emergence of variants which exhibit increased growth rates compared to the intact adenovector. With continued passaging of the original Ad gag vector, the level of variants increases and hence amplification rates increase also.

The MRK Ad5 HIV-1 gag virus has also been continually passaged under process conditions (i.e., serum-free media). Viral DNA extracted from passages 11 and 12 show no evidence of rearrangement.

Table 4:
Amplification Ratios Based on AEX and QPA Analysis of
Virus Amplification from Passage 3 to Passage 4.

Ad gag construct	Amplification Ratio
MRKAd5gag	470
HCMV-Flgag-bGHpA [E3-]	115
HCMV-Flgag-SPA [E3+]	320
mCMV-FLgag-bGHpA [E3+]	420
Original construct *	40 - 50

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EXAMPLE 13

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Analytical Evaluation of the enhanced Ad5 Constructs

To study the effects of the transgene and the E3 gene on virus amplification,
the enhanced adenoviral vector, MRK Ad5 HIV-1 gag, along with its transgene-less

version (MRKpAdHVE3) and its E3- version (MRK Ad5 HIV-1 gag E3-), was studied for several passages under serum-free conditions. Table 5A shows the amplification ratios determined for passages P3 to P8 for MRK Ad5 HIV-1 gag. Within a certain MOI range, it has been determined that the virus output is directly proportional to the virus input. Therefore, the greater the number of virus particles

per cell at infection, the greater the virus amount produced. Viral amplification ratios, on the other hand, are inversely proportional to the virus input. The lower the virus input, the greater the amplification ratio.

Table 5B shows the amplification rates of the new E3+ vector backbone MRKpAdHVE3. It has a significantly lower rate of amplification compared with the gag transgene containing version. This may be contributed to the larger size MRK Ad5 HIV-1 gag since it contains the transgene. This inclusion of the transgene brings the size of the adenovirus closer to the size of a wild type Ad5 virus. It is well known that adenoviruses amplify best when they are at close to their wild type genomic size.

^{*} This estimation is based on the clinical lot growth characteristics at Passage 12.

Wild type Ad5 is 35,935 bp. The MRKpAdHVE3 is 32, 905 bp in length. The enhanced adenovector MRK Ad5 HIV-1 gag is 35,453bp (See Figure 14 for vector map; see also Figure 15A-X show the complete pre-adenoviral vector sequence, which includes an additional 2,021 bp of the vector backbone).

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Table 5C shows the amplification rates of the new E3- gag containing virus MRK Ad5 HIV-1 gag E3-. Once again, this virus shows lower growth rate than the enhanced adenoviral vector. This may be attributed to the decreased sized of this virus (due to the E3 gene deletion) compared with wild type Ad5. The MRK Ad5 HIV-1 gag E3- virus is 32,810 bp in length. This can be compared with the wild type Ad5 which is 35,935 bp and MRK Ad5 HIV-1 gag which is 35,453 bp in length.

Table 5A: Amplification ratios determined by AEX and QPA for MRKAd5gag over several continuous passaging in serum free media. Following P5, two replicate samples were taken (rep-1 and rep-2) and analyzed.

MRKAd5gag rep1

	XV (10° calls/tr	D, Vlability (%)	Harvest Time	Cell Passage	Titor	TRer	OPA	Ratio	Amplification	AEX
	Infaction	Harvest	hpl	Number	10 rd vp/ml cultura	10° vp/call	10° TCID _{so} /ml	AEX:OPA	Ratio	Internal Control
P4	1,49, 81%	0.58, 50%	44	46	8.7	5.9	1.72	50	470 (MOI = 125)	
P5	1.38, 93%	0.66, 47%	48	49	6.7	4.9	1.38	49	170	
P6	1,04, 94%	0.68, 77%	47	48	5.8	5.6	1.42	41	200	
P7	1.50, 84%	0.96, 61%	49.5	50	3.9	1.4	0.87	40	50	
P7	1.09, 97%	0.76, 59%	50	52	5.2	4.7	1.70	81	170	
P8	1.03, 94%	0.85, 64%	47.5	54	9.0	8.7	1.10	82	310	
P9	0.89, 95%	0.99, 73%	47.5	56	4,4	4.9	1.03	43	175	3.12 2.84
P10	1.09, 91%	1.06, 86%	47.5	58	8.0	2.8	1.18	26	100	2.70 2.60
P11	1.19, 88%	0.88, 65%	47	60	3.6	3.0	1.15	31	110	2.70 2.70
P12	0.58, 91%	0.85, 63%	47.5	47	5.4	5.5	1.20	45	200	2.86 2.60
P13	1.00, 88%	0.70, 67%	49	49	5.8	5.8	1.11	52	210	3.18 3.18
P14	1.94, 92%	0.88, 67%	46	53	8.6	4.4			160	3.28 3.27
P15	0.97, 96%	0.64, 66%	47	47	6.9	7.1			250	3.12 2.91

Table 5B: Amplification ratios determined by AEX and QPA for MRKHVE3 over several continuous passaging in serum free media. MRKHVE3 is the new vector backbone which does NOT carry a transgene.

MRKHVE3

	Xy (10° cells/m	I), Viability (%)	Harvest Time	Cell Passage	Titer	Titer	QPA .	Ratio	Amplification	AEX
	Infection	Harvest	h.p.l.	Number	10 ^{to} vp/mi culture	10° vp/ceti	10° TCID _{to} /ml	AEX:QPA	Ratio	Internal Control
P4	1.10, 97%	1.28, 79%	49	54	4.1	3.8	1.70	25	300 (MC) = 125)	
P5	0.92, 89%	1.18, 77%	47	. 48	4.3	4.7	1.24	35	170	
P6	1.55, 88%	1.26, 76%	49.5	50	1.2	0.8	0.58	21	30	
P6	1.09, 97%	1.11, 81%	49	52	4.0	3.6	1.18	34	130	
P7	1.17, 91%	1.22, 91%	47.5	54	3.7	3.2	0.50	74	110	
P8	0.98, 88%	1.41, 83%	48	56	2.1	2.1	0,47	45	75	3.12 2.84
P9	1.20, 89%	1.25, 81%	47.5	58	8.0	0.7	0,29	28	25	2.70 2.60
P10	0.99, 82%	1.55, 88%	47	60	2.3	2.3	0.43	53	80	2.70 2.70
P11	1.07, 96%	1.25, 83%	48	47	2.7	2.5	0.41	66	90	2.86 2.60
P12	0.80, 91%	1.14, 80%	49.5	49	6.9	7.4	0.48	123	260	3.18 3.18
P13	1.96, 95%	1.14, 85%	45.5	53	5.8	3.0			110	3.28 3.27
P14	0.97, 96%	1.03, 98%	48.5	47	9.4	8.7			350	3.12 2.91
P15	0.87, 99%	0.97, 59%	49.5	49	5.3	6.1	†		218	2.78 2.52

Table 5C. Amplification ratios determined by AEX and QPA for MRKAd5gag(E3-) over several continuous passaging in serum free media. This construct is identical to the MRKAd5gag construct except that this version is DELETED of the E3 gene.

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MRKAd5gag(E3-)

		il), Viability (%)	Harvest Time	Cell Passage	Titler	Titer	QPA	Ratio	Amplification	AEX
	Infection	Harvest	h.p.L	Number	10 ⁴⁰ vp/mi culture	10° vp/cell	10° TCID _{EP} Init	AEX:QPA	Ratio	Internal Control
P4	1.62, 77%	1.12, 62%	47.5	46	2.0	1.2	0.92	20	100 (MOI=125)	
P5	1.16, 92%	0.62, 43%	49	49	3.3	2.9	0.99	34	100	
P6	1.71, 86%	0.20, 10%	49	50	4.7	2.7	1.70	28	100	
P6	1.09, 97%	0.63, 54%	49.5	52	5.4	5.0	1.76	31	180	
P7	1.17, 91%	0.98, 72%	47.50	54	7.1	6.1	0.67	106	220	
P8	0.98, 88%	0.77, 48%	48	56	3.1	32	0.66	47	115	3.12 2.84
P9	1.20, 89%	1.03, 72%	48	58	1.8	1.5	0.57	32	55	2.70 2.60
P10	0.99, 82%	0.80, 62%	46.5	60	3.2	3.2	83.0	47	115	2.70 2.70
P11	1.07, 96%	0.88, 70%	48.5	47	5.9	5.5	83.0	87	200	2.88 2.60
P12	0.80, 91%	0.67, 59%	50	49	5.1	6.4	0.72	71	230	3.18 3.18
P13	1.96, 95%	0.91, 59%	45.5	53	7.4	3.8		l	135	3.28 3.27
P14	0.97, 96%	0.81, 74%	48	47	6.8	7.0			250	3.12 2.91
P15	0.87, 99%	0.84, 56%	49	49	4.8	5.5			196	2.78 2.52

EXAMPLE 14

Gag Expression Analysis of the Novel Constructs

In vitro gag analysis of the MRK Ad5 HIV-1 gag and the original HIV-gag vectors (research and clinical lot) show comparable gag expression. The clinical lot shows only a slightly reduced gag expression level. The most noticeable difference is with the mCMV vector. This vector shows roughly 3 fold lower expression levels compared with the other vectors tested (which all contain hCMV promoters). The mCMV-FLgag with bGHpA assay was performed three times using different propagation and purification lots and it consistently exhibited weaker gag expression.

EXAMPLE 15

Evaluation of MRK Ad5 HIV-1 gag and Other gag-Containing Adenovectors in Balb/c Mice

Cohorts of 10 balb/c mice were vaccinated intramuscularly with escalating doses of MRK Ad5 HIV-1 gag, and the research and clinical lots of original Ad5HIV-1gag. Serum samples were collected 3 weeks post dose 1 and analyzed by anti-p24 sandwich ELISA.

Anti-p24 titers in mice that received MRK Ad5 HIV-1 gag (107 and 109 vp(viral particle) doses) were comparable (Figure 13) to those of the research lot of Ad5HIV-1 gag, for which much of the early rhesus data were generated on. These titers were also comparable when E3 is deleted (MRKAd5hCMVgagbGHpA(E3-)) or SPA is substituted for bGHpA terminator (MRKAd5 hCMV-gag-SPA (E3+)) or murine CMV promoter is used in place of hCMV (MRKAd5 mCMV-gag-bGHpA (E3+)) in the MRKAd5 backbone.

The results shown in Table 7 indicate that the three other vectors (in addition to the preferred vector, MRK Ad5 HIV-1 gag, are also capable of inducing strong anti-gag antibody responses in mice. Interestingly enough, while the mCMV-FLgag construct containing bGHpA and E3+ in an E1 parallel orientation showed lowest gag expression in the COS cell *in vitro* infection (Table 6) in comparison with the other vectors tested, it generated the greatest anti-gag antibody response this *in vivo* Balb/c study. Table 7 also shows a dose response in anti-gag antibody production in both the research and the clinical lot. As expected, the clinical lot shows reduced anti-gag antibody induction at each dosage level compared to the same dosage used for the research lot.

Table 6: In vitro analysis for gag expression in COS cells by Elisa assay.

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Viral Vectors ^a	μg gag/4.8x10e5 COS/10e8 parts/48hr
MRKAd5gag ^b	1.40
Clinical lot Ad5gag ^c	1.28
Research lot Ad5gag ^d	1.32
MCMVFL-gagbGHpA ^e	0.42

^a A_{260am} absorbance readings taken for viral particle determinations.

^b MRKAd5gag was produced in serum free conditions and purified at P5.

^c Clinical lot# Ad5gagFN0001

²⁵ dResearch Ad5FLgag lot# 6399

^e mCMVFL-gagbGHpA was produced in serum free conditions and purified at P5.

Table 7: mHIV020 Anti-p24 Ab Titers in Balb/c mice (n=10) vaccinated with various Adgag constructs and lots (3 week post dose1).

Group ID	Vaccine	Dose (vp)	GMT	SE upper	SE lower
1	^a MRKAd5gag	10^7	25600	5877	4780
2	в	10^9	409600	94028	76473
3	hCMV FL-gag bGHpA [E3-] →	10^7	7352	2077	1620
4	g .	10^9	235253	59767	47659
5	hCMV FL-gag SPA [E3+] →	10^7	12800	9905	236
6	•	10^9	310419	99181	75165
7	^b mCMV FL-gag bGHpA [E3+] →	10^7	44572	23504	15389
8	-	10^9	941014	239068	190636
9	^c hCMV FL-gag bGHpA [E3-] ←	10^7	3676	934	745
10		10^9	117627	17491	15227
11	research lot hCMV intronA FL-gag bGHpA [E3-] <-	10^6 10^7	528 14703	262 5274	175 3882
12	и	10/8	58813	14942	11915
14	*	10^9	204800	53232	42250
15	clinical lot hCMVintronA FL-gag bGHpA [E3-] <-	10^6	230	82	61
16 17		10^7	4222 19401	3405 3939	1138 3274
18	n	10^9	89144	25187	19639
19	Naïve	none	93	7	6

*2x50 µL i.m. (quad) injections/animal P.I.s: Youil, Chen, Casimiro

Vaccination: T. Toner, Q. Su

Assay: M. Chen

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^aThe structure of MRKAd5gag is: hCMVFL-gagbGHpA [E3+] → The <u>same lot</u> of MRKAd5gag used in this rodent study was used in the Rhesus monkey study (Tables 7 and 8).

^bThe same lot of mCMVFL-gagbGHpA[E3+] used in the *in vitro* study (Table 6) ws used here.

^cThis construct was designed by Volker Sandig. It contains a shorter version of the hCMV

This construct was designed by Volker Sandig. It contains a shorter version of the hCMV promoter than that used in the MRK constructs. The adenovector backbone is identical to the original backbone used in the original Adgag vector. Expression at 10e7 dose from this vector is 7 fold lower then the same dose of the MRKAd5gag and 4 fold lower than the research lot.

EXAMPLE 16

Comparison of Humoral and Cellular Responses Towards the Original Ad-gag Construct with the New MRK Ad5 HIV-1 gag in Rhesus Monkeys

Cohorts of 3 rhesus monkeys were vaccinated intramuscularly with MRK Ad5 HIV-1 gag or the clinical Ad5gag bulk at two doses, 10^{11} vp and 10^9 vp. Immunizations were conducted at week 0, 4, and 25. Serum and PBMC samples were collected at selected time points. The serum sample were assayed for anti-p24 Ab titers (using competitive based assay) and the PBMCs for antigen-specific IFN-gamma secretion following overnight stimulation with gag 20-mer peptide pool (via ELISpot assay).

The results shown in Table 8 indicate comparable responses with respect to the generation of anti-gag antibodies. The frequencies of gag-specific T cells in

peripheral blood assummarized in Table 9 demonstrate a strong cellular immune response generated after a single dose with the new construct MRK Ad5 HIV-1 gag. The responses are also boostable with second dose of the same vector. The vector is also able to induce CD8+ T cell responses (as evident by remaining spot counts after CD4+ depletion of PBMCs) which are responsible for cytotoxic activity.

Table 8 Anti-p24 antibody titers (in mMU/mL) in rhesus macaques immunized with

gag-expressing adenovectors (Protocol HIV203).

yag-expressing adenovecu	Pre	Wk4	Wk 8	Wk 12	Wk 16	Wk 20	Wk 25	Wk 28
MRKAd5gag ^a , 10^11 vp								
97N010	<10	118	5528	11523	7062	21997	ND	51593
97N116	<10	62	772	1447	1562	2174	ND	20029
98X007	<10	66.	3353	6156	6845	3719	ND_	24031
MR K Ad5gag, 10^9 vp								
97N120	<10_	51	204	318	366	482	ND	6550
97N144	<10	18	118_	274	706	888	ND	7136
98X008	<10	15	444	386	996_	1072	ND_	12851
Ad5gog ^b , Clinical Lot, 10^11 vp	† — —							
97X001	<10	87	2579	4718	7174	7250	ND	69226
97N146	<10	72	3604	7380	7526	18906	ND	60283
98X009	<10	78	4183	3946	3124	6956_	ND_	26226
Ad5gag, Clinical Lat, 10^9 vp			<u> </u>	<u> </u>				
97N020	<10	<10	143_	371	390	1821	ND	17177
97X003	<10	<10	39	93	156	596_	ND	2053
98X012	<10	81	342	717	956	1558_	ND	11861
MRKAdagag (hCMV, bGHpA, E3+)	1			1	 		
bariginal Actigag vector (hCMV/Intr	on A both	A, E3-), lot	#FN0001	<u> </u>		 	 	
ND, not determined			<u> </u>	<u> </u>	<u> </u>			1

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Table 9. Number of gag-specific T cells per million peripheral blood mononuclear cells (PBMCs) in rhesus monkeys immunized with gag-expressing adenovectors.

Also included are those frequencies in PBMCs depleted of CD4⁺ T cells.

Grp #	Vaccination	Monkey ID	T=4	Wk	T=6	Wk	1=1	Wk	I=16	5 Wk	T #2	. Wk	T=28	Wk
	7=0.4.25 wks		Media	Gog H ^b	Media	Gog H	Media	Gog H	Media	Gog H	Media	Gog H	Media	Gog H
		97ND1D	6	89	0	395	0	1058	0	1174	3	775	4	1074
]]	MRKAGOOD	97N010(CD4-)	۱۵	38	"	0.55	3	993	•	11/4	ا ة ا	76	lõi	594
1	10^11 Vp	97N116	l i	398	1 1	609	ŏ	534	4	395	lĭ	261	lõ	408
į.		97N116(CD4-)	ii	676	! '	-	ŏ	593		","	Ó	184	ŏ	666
ì		98X007	10	579	0	1304	1 3	2193	ı	2118	l 3	1588	ŏ	2113
ł		98X007(CD4-)	20	965	ľ		ŏ	2675	i '		Ιŏ	1656	ا ة ا	1278
		YOUU/(CLAF)		700	ĺ	1	ı .	20/0			Ĭ		L'	
2	MRKAdaga	97N120	5	275	1	249	4	141	4	119	9	206	4	219
<u> </u>	10/9 VP	97N120(CD4-)	11	170	1		0	85			0	75	1 1	219
1	-	97N144	3	235	6	438	וון	318	3	256	1 1	98	5	373
		97N144(CD4-)	6	148	1		0	285			ND	ND	0	625
		98X008	4	368	1	1090	3	891	4	673	3	473	5	735
}		98X008(CD4-)	14	696	1	1	0	1175	1	1	0	391	4	848
		077/001	 0	261	1	485	-	817	0	12200	- , -	894	0	1858
3 .	AcSgoop clinical lat	97X001		283	1 .	465	3	996	I۳	12000	اة ا	1010	Ιŏ	1123
1	10^11 vp	97X001(CD4-) 97N146	10	150	1	465	١٥	339	١,	1272	3	1238	l š	1785
1			1	133	1 '	~~~	lő	370	i '	1272	ŏ	654	۱ŏ	971
1		97N146(CD4-) 98X009	6	93	Ìз	339	1 3	559	0	896	1 1	384	lŏ	1748
1			۱ŏ	73	1 "		۱۵	333	١٠	""	1 6	225	0	644
		98X009(CD4-)	١ '	/3	1	1	ľ	333			ľ	1 23	L	1
4	Adsaca dinical lat	97N020	3	30	1	101	0	66	0	36	0	26	0	41
	10/9 vp	97N020(CD4-)	10	29	Į .	l	0	15	١.	١	l o	1 1	0	16
Į.		97X003	4	68	5	134	0	18	יו	38	4	38	6	81
1		97X003(CD4-)	9	40	1	•	0	٥	1	1	0	4	0	19
I		98X012	5	95	3	54	1 1	34	0	18	0	20	1 1	121
		98X012(CD4-)	11	70		ĺ	0	111		1	0	В	0	41
5	Nave	96R041	6	8	17	1.	0	0	0	0	0	0	1	0
		053F	14	18	5	16.	20	14	19	15	10	15	24	١ ٠
1			1											

Based on either 4x10/6 or 2x10/5 cells per well (depending on spot density)

ND, not determined

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"mock or no peptide control

The adenovectors described herein and, particularly, MRK Ad5 HIV-1 gag, represent very promising HIV-gag adenovectors with respect to their enhanced growth characteristics in both serum and, more importantly, in serum-free media conditions. In comparison with the current HIV-1 gag adenovector construct, MRK Ad5 HIV-1 gag shows a 5-10 fold increased amplification rate. We have shown that it is genetically stable at passage 21. This construct is able to generate significant cellular immune responses in vivo even at a relatively low dose of 10^9 vp. The potency of the MRKAd5gag construct is comparable to, if not better than the original HIV-1gag vector as shown in this rhesus monkey study.

EXAMPLE 17 CODON OPTIMIZED HIV-1 POL AND CODON OPTIMZED HIV-1 POL MODIFICATIONS

The open reading frames for the various synthetic *pol* genes disclosed herein comprise coding sequences for the reverse transcriptase (or RT which consists of a polymerase and RNase H activity) and integrase (IN). The protein sequence is based

Pod of 20-capeptides overlapping by 10 accord encompassing the gas sequence

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on that of Hxb2r, a clonal isolate of IIIB; this sequence has been shown to be closest to the consensus clade B sequence with only 16 nonidentical residues out of 848 (Korber, et al., 1998, Human retroviruses and AIDS, Los Alamos National Laboratory, Los Alamos, New Mexico). The skilled artisan will understand after review of this specification that any available HIV-1 or HIV-2 strain provides a potential template for the generation of HIV pol DNA vaccine constructs disclosed herein. It is further noted that the protease gene is excluded from the DNA vaccine constructs of the present invention to insure safety from any residual protease activity in spite of mutational inactivation. The design of the gene sequences for both wildtype (wt-pol) and inactivated pol (IA-pol) incorporates the use of human preferred ("humanized") codons for each amino acid residue in the sequence in order to maximize in vivo mammalian expression (Lathe, 1985, J. Mol. Biol. 183:1-12). As can be discerned by inspecting the codon usage in SEQ ID NOs: 1, 3, 5 and 7, the following codon usage for mammalian optimization is preferred: Met (ATG), Gly (GGC), Lys (AAG), Trp (TGG), Ser (TCC), Arg (AGG), Val (GTG), Pro (CCC), Thr (ACC), Glu (GAG); Leu (CTG), His (CAC), Ile (ATC), Asn (AAC), Cys (TGC), Ala (GCC), Gln (CAG), Phe (TTC) and Tyr (TAC). For an additional discussion relating to mammalian (human) codon optimization, see WO 97/31115 (PCT/US97/02294), which, as noted elsewhere in this specification, is hereby incorporated by reference. It is intended that the skilled artisan may use alternative versions of codon optimization or may omit this step when generating HIV pol vaccine constructs within the scope of the present invention. Therefore, the present invention also relates to non-codon optimized versions of DNA molecules and associated recombinant adenoviral HIV vaccines which encode the various wild type and modified forms of the HIV Pol protein disclosed herein. However, codon optimization of these constructs is a 25 preferred embodiment of this invention.

A particular embodiment of this portion of the invention comprisies codon optimized nucleotide sequences which encode wt-pol DNA constructs (herein, "wtpol" or "wt-pol (codon optimized))" wherein DNA sequences encoding the protease (PR) activity are deleted, leaving codon optimized "wild type" sequences which encode RT (reverse transcriptase and RNase H activity) and IN integrase activity. A DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:1, the open reading frame being contained from an initiating Met residue at nucleotides 10-12 to a termination codon from nucleotides 2560-2562. SEQ ID NO:1 is as follows: AGATCTACCA TGGCCCCCAT CTCCCCCATT GAGACTGTGC CTGTGAAGCT GAAGCCTGGC ATGGATGGCC CCAAGGTGAA GCAGTGGCCC CTGACTGAGG AGAAGATCAA GGCCCTGGTG

	GAAATCTGCA	CTGAGATGGA	GAAGGAGGC	AAAATCTCCA	AGATTGGCCC	CGAGAACCCC
	TACAACACCC	CTGTGTTTGC	CATCAAGAAG	AAGGACTCCA	CCAAGTGGAG	GAAGCTGGTG
	GACTTCAGGG	AGCTGAACAA	GAGGACCCAG	GACTTCTGGG	AGGTGCAGCT	GGGCATCCCC
	CACCCCGCTG	GCCTGAAGAA	GAAGAAGTCT	GTGACTGTGC	TGGATGTGGG	GGATGCCTAC
5	TTCTCTGTGC	CCCTGGATGA	GGACTTCAGG	AAGTACACTG	CCTTCACCAT	CCCCTCCATC
	AACAATGAGA	CCCCTGGCAT	CAGGTACCAG	TACAATGTGC	TGCCCCAGGG	CTGGAAGGGC
	TCCCCTGCCA	TCTTCCAGTC	CTCCATGACC	AAGATCCTGG	AGCCCTTCAG	GAAGCAGAAC
	CCTGACATTG	TGATCTACCA	GTACATGGAT	GACCTGTATG	TGGGCTCTGA	CCTGGAGATT
	GGGCAGCACA	GGACCAAGAT	TGAGGAGCTG	AGGCAGCACC	TGCTGAGGTG	GGGCCTGACC
10	ACCCCTGACA	AGAAGCACCA	GAAGGAGCCC	CCCTTCCTGT	GGATGGGCTA	TGAGCTGCAC
	CCCGACAAGT	GGACTGTGCA	GCCCATTGTG	CTGCCTGAGA	AGGACTCCTG	GACTGTGAAT
	GACATCCAGA	AGCTGGTGGG	CAAGCTGAAC	TGGGCCTCCC	AAATCTACCC	TGGCATCAAG
	GTGAGGCAGC	TGTGCAAGCT	GCTGAGGGGC	ACCAAGGCCC	TGACTGAGGT	GATCCCCCTG
	ACTGAGGAGG	CTGAGCTGGA	GCTGGCTGAG	AACAGGGAGA	TCCTGAAGGA	GCCTGTGCAT
15	GGGGTGTACT	ATGACCCCTC	CAAGGACCTG	ATTGCTGAGA	TCCAGAAGCA	GGGCCAGGGC
	CAGTGGACCT	ACCAAATCTA	CCAGGAGCCC	TTCAAGAACC	TGAAGACTGG	CAAGTATGCC
	AGGATGAGGG	GGGCCCACAC	CAATGATGTG	AAGCAGCTGA	CTGAGGCTGT	GCAGAAGATC
	ACCACTGAGT	CCATTGTGAT	CTGGGGCAAG	ACCCCCAAGT	TCAAGCTGCC	CATCCAGAAG
	GAGACCTGGG	AGACCTGGTG	GACTGAGTAC	TGGCAGGCCA	CCTGGATCCC	TGAGTGGGAG
20	TTTGTGAACA	CCCCCCCT	GGTGAAGCTG	TGGTACCAGC	TGGAGAAGGA	GCCCATTGTG
	GGGGCTGAGA	CCTTCTATGT	GGATGGGGCT	GCCAACAGGG	AGACCAAGCT	GGGCAAGGCT
	GGCTATGTGA	CCAACAGGGG	CAGGCAGAAG	GTGGTGACCC	TGACTGACAC	CACCAACCAG
	AAGACTGAGC	TCCAGGCCAT	CTACCTGGCC	CTCCAGGACT	CTGGCCTGGA	GGTGAACATT
	GTGACTGACT	CCCAGTATGC	CCTGGGCATC	ATCCAGGCCC	AGCCTGATCA	GTCTGAGTCT
25	GAGCTGGTGA	ACCAGATCAT	TGAGCAGCTG	ATCAAGAAGG	AGAAGGTGTA	CCTGGCCTGG
	GTGCCTGCCC	ACAAGGGCAT	TGGGGGCAAT	GAGCAGGTGG	ACAAGCTGGT	GTCTGCTGGC
	ATCAGGAAGG	TGCTGTTCCT	GGATGGCATT	GACAAGGCCC	AGGATGAGCA	TGAGAAGTAC
	CACTCCAACT	GGAGGGCTAT	GGCCTCTGAC	TTCAACCTGC	CCCCTGTGGT	GGCTAAGGAG
	ATTGTGGCCT	CCTGTGACAA	GTGCCAGCT	AAGGGGGAGG	CCATGCATGG	GCAGGTGGAC
30	TGCTCCCCTG	GCATCTGGC	GCTGGACTG	ACCCACCTGG	AGGGCAAGGT	GATCCTGGTG
	GCTGTGCATG	TGGCCTCCGG	CTACATTGA	GCTGAGGTGA	1 TCCCTGCTGA	GACAGGCCAG
	GAGACTGCCT	ACTTCCTGCT	GAAGCTGGC	r ggcaggtgg(CTGTGAAGAC	CATCCACACT
	GACAATGGCT	CCAACTTCAC	TGGGGCCAC	A GTGAGGGCT	CCTGCTGGTG	GGCTGGCATC
	AAGCAGGAGT	TTGGCATCC	CTACAACCC	CAGTCCCAG	GGGTGGTGG	GTCCATGAAC
35	AAGGAGCTG	AGAAGATCA	TGGGCAGGT	G AGGGACCAG	CTGAGCACC	GAAGACAGCT
	GTGCAGATG	CTGTGTTCA	CCACAACTT	C AAGAGGAAG	G GGGGCATCG	GGGCTACTCC

GCTGGGGAGA GGATTGTGGA CATCATTGCC ACAGACATCC AGACCAAGGA GCTCCAGAAG
CAGATCACCA AGATCCAGAA CTTCAGGGTG TACTACAGGG ACTCCAGGAA CCCCCTGTGG
AAGGGCCCTG CCAAGCTGCT GTGGAAGGGG GAGGGGGCTG TGGTGATCCA GGACAACTCT
GACATCAAGG TGGTGCCCAG GAGGAAGGCC AAGATCATCA GGGACTATGG CAAGCAGATG
GCTGGGGATG ACTGTGTGGC CTCCAGGCAG GATGAGGACT AAAGCCCGGG CAGATCT (SEQ
ID NO:1).

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The open reading frame of the wild type pol construct disclosed as SEQ ID NO:1 contains 850 amino acids, disclosed herein as SEQ ID NO:2, as follows: Met Ala Pro Ile Ser Pro Ile Glu Thr Val Pro Val Lys Leu Lys Pro Gly Met Asp Gly Pro Lys Val Lys Gln Trp Pro Leu Thr Glu Glu Lys 10 Ile Lys Ala Leu Val Glu Ile Cys Thr Glu Met Glu Lys Glu Gly Lys Ile Ser Lys Ile Gly Pro Glu Asn Pro Tyr Asn Thr Pro Val Phe Ala Ile Lys Lys Lys Asp Ser Thr Lys Trp Arg Lys Leu Val Asp Phe Arg Glu Leu Asn Lys Arg Thr Gln Asp Phe Trp Glu Val Gln Leu Gly Ile 15 Pro His Pro Ala Gly Leu Lys Lys Lys Ser Val Thr Val Leu Asp Val Gly Asp Ala Tyr Phe Ser Val Pro Leu Asp Glu Asp Phe Arg Lys Tyr Thr Ala Phe Thr Ile Pro Ser Ile Asn Asn Glu Thr Pro Gly Ile Arg Tyr Gln Tyr Asn Val Leu Pro Gln Gly Trp Lys Gly Ser Pro Ala Ile Phe Gln Ser Ser Met Thr Lys Ile Leu Glu Pro Phe Arg Lys Gln Asn Pro Asp Ile Val Ile Tyr Gln Tyr Met Asp Asp Leu Tyr Val Gly 20 Ser Asp Leu Glu Ile Gly Gln His Arg Thr Lys Ile Glu Glu Leu Arg Gln His Leu Leu Arg Trp Gly Leu Thr Thr Pro Asp Lys Lys His Gln Lys Glu Pro Pro Phe Leu Trp Met Gly Tyr Glu Leu His Pro Asp Lys Trp Thr Val Gln Pro Ile Val Leu Pro Glu Lys Asp Ser Trp Thr Val 25 Asn Asp Ile Gln Lys Leu Val Gly Lys Leu Asn Trp Ala Ser Gln Ile Tyr Pro Gly Ile Lys Val Arg Gln Leu Cys Lys Leu Leu Arg Gly Thr Lys Ala Leu Thr Glu Val Ile Pro Leu Thr Glu Glu Ala Glu Leu Glu Leu Ala Glu Asn Arg Glu Ile Leu Lys Glu Pro Val His Gly Val Tyr Tyr Asp Pro Ser Lys Asp Leu Ile Ala Glu Ile Gln Lys Gln Gly Gln 30 Gly Gln Trp Thr Tyr Gln Ile Tyr Gln Glu Pro Phe Lys Asn Leu Lys Thr Gly Lys Tyr Ala Arg Met Arg Gly Ala His Thr Asn Asp Val Lys Gln Leu Thr Glu Ala Val Gln Lys Ile Thr Thr Glu Ser Ile Val Ile Trp Gly Lys Thr Pro Lys Phe Lys Leu Pro Ile Gln Lys Glu Thr Trp Glu Thr Trp Trp Thr Glu Tyr Trp Gln Ala Thr Trp Ile Pro Glu Trp 35 Glu Phe Val Asn Thr Pro Pro Leu Val Lys Leu Trp Tyr Gln Leu Glu Lys Glu Pro Ile Val Gly Ala Glu Thr Phe Tyr Val Asp Gly Ala Ala

Asn Arg Glu Thr Lys Leu Gly Lys Ala Gly Tyr Val Thr Asn Arg Gly Arg Gln Lys Val Val Thr Leu Thr Asp Thr Thr Asn Gln Lys Thr Glu Leu Gln Ala Ile Tyr Leu Ala Leu Gln Asp Ser Gly Leu Glu Val Asn Ile Val Thr Asp Ser Gln Tyr Ala Leu Gly Ile Ile Gln Ala Gln Pro Asp Gln Ser Glu Ser Glu Leu Val Asn Gln Ile Ile Glu Gln Leu Ile 5 Lys Lys Glu Lys Val Tyr Leu Ala Trp Val Pro Ala His Lys Gly Ile Gly Gly Asn Glu Gln Val Asp Lys Leu Val Ser Ala Gly Ile Arg Lys Val Leu Phe Leu Asp Gly Ile Asp Lys Ala Gln Asp Glu His Glu Lys Tvr His Ser Asn Trp Arg Ala Met Ala Ser Asp Phe Asn Leu Pro Pro Val Val Ala Lys Glu Ile Val Ala Ser Cys Asp Lys Cys Gln Leu Lys 10 Gly Glu Ala Met His Gly Gln Val Asp Cys Ser Pro Gly Ile Trp Gln Leu Asp Cys Thr His Leu Glu Gly Lys Val Ile Leu Val Ala Val His Val Ala Ser Gly Tyr Ile Glu Ala Glu Val Ile Pro Ala Glu Thr Gly Gln Glu Thr Ala Tyr Phe Leu Leu Lys Leu Ala Gly Arg Trp Pro Val Lys Thr Ile His Thr Asp Asn Gly Ser Asn Phe Thr Gly Ala Thr Val 15 Arg Ala Ala Cys Trp Trp Ala Gly Ile Lys Gln Glu Phe Gly Ile Pro Tyr Asn Pro Gln Ser Gln Gly Val Val Glu Ser Met Asn Lys Glu Leu Lys Lys Ile Ile Gly Gln Val Arg Asp Gln Ala Glu His Leu Lys Thr Ala Val Gln Met Ala Val Phe Ile His Asn Phe Lys Arg Lys Gly Gly 20 Ile Gly Gly Tyr Ser Ala Gly Glu Arg Ile Val Asp Ile Ile Ala Thr Asp Ile Gln Thr Lys Glu Leu Gln Lys Gln Ile Thr Lys Ile Gln Asn Phe Arg Val Tyr Tyr Arg Asp Ser Arg Asn Pro Leu Trp Lys Gly Pro Ala Lys Leu Trp Lys Gly Glu Gly Ala Val Val Ile Gln Asp Asn Ser Asp Ile Lys Val Val Pro Arg Arg Lys Ala Lys Ile Ile Arg Asp 25 Tyr Gly Lys Gln Met Ala Gly Asp Asp Cys Val Ala Ser Arg Gln Asp Glu Asp (SEQ ID NO:2).

The present invention especially relates to an adenoviral vector vaccine which comprises a codon optimized HIV-1 DNA pol construct wherein, in addition to deletion of the portion of the wild type sequence encoding the protease activity, a combination of active site residue mutations are introduced which are deleterious to HIV-1 pol (RT-RH-IN) activity of the expressed protein. Therefore, the present invention preferably relates to an adenoviral HIV-1 DNA pol-based vaccine wherein the construct is devoid of DNA sequences encoding any PR activity, as well as containing a mutation(s) which at least partially, and preferably substantially, abolishes RT, RNase and/or IN activity. One type of HIV-1 pol mutant which is part and parcel of an adenoviral vector vaccine may include but is not limited to a mutated

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DNA molecule comprising at least one nucleotide substitution which results in a point mutation which effectively alters an active site within the RT, RNase and/or IN regions of the expressed protein, resulting in at least substantially decreased enzymatic activity for the RT, RNase H and/or IN functions of HIV-1 Pol. In a preferred embodiment of this portion of the invention, a HIV-1 DNA pol construct contains a mutation or mutations within the Pol coding region which effectively abolishes RT, RNase H and IN activity. An especially preferable HIV-1 DNA pol construct in a DNA molecule which contains at least one point mutation which alters the active site of the RT, RNase H and IN domains of Pol, such that each activity is at least substantially abolished. Such a HIV-1 Pol mutant will most likely comprise at least one point mutation in or around each catalytic domain responsible for RT, RNase H and IN activity, respectfully. To this end, an especially preferred HIV-1 DNA pol construct is exemplified herein and contains nine codon substitution mutations which results in an inactivated Pol protein (IA Pol: SEQ ID NO:4, Figure 17A-C) which has no PR, RT, RNase or IN activity, wherein three such point mutations reside within each of the RT, RNase and IN catalytic domains. Therefore, an especially preferred exemplification is an adenoviral vaccine which comprises, in an appropriate fashion, a DNA molecule which encodes IA-pol, which contains all nine mutations as shown below in Table 1. An additional preferred amino acid residue for substitution is Asp551, localized within the RNase domain of Pol. Any combination of the mutations disclosed herein may suitable and therefore may be utilized as an IA-Pol-based vaccine of the present invention. While addition and deletion mutations are contemplated and within the scope of the invention, the preferred mutation is a point mutation resulting in a substitution of the wild type amino acid with an alternative amino acid residue.

Т	'a	bl	le	1

	wt aa	aa residue	mutant aa	enzyme function
	Asp	112	Ala	RT
	Asp	187	Ala	RT
30	Asp	188	Ala	RT
	Asp .	445	. Ala	. RNase H
	Glu	480	Ala	RNase H
	Asp	500	Ala	RNase H
	Asp	626	Ala	IN
	Asp	678	Ala	IN
	Glu	714	Ala	IN

It is preferred that point mutations be incorporated into the IApol mutant adenoviral vaccines of the present invention so as to lessen the possibility of altering epitopes in and around the active site(s) of HIV-1 Pol.

To this end, SEQ ID NO:3 discloses the nucleotide sequence which codes for a codon optimized pol in addition to the nine mutations shown in Table 1, disclosed as follows, and referred to herein as "IApol":

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AGATCTACCA TGGCCCCCAT CTCCCCCATT GAGACTGTGC CTGTGAAGCT GAAGCCTGGC ATGGATGGCC CCAAGGTGAA GCAGTGGCCC CTGACTGAGG AGAAGATCAA GGCCCTGGTG GAAATCTGCA CTGAGATGGA GAAGGAGGGC AAAATCTCCA AGATTGGCCC CGAGAACCCC TACAACACCC CTGTGTTTGC CATCAAGAAG AAGGACTCCA CCAAGTGGAG GAAGCTGGTG 10 GACTTCAGGG AGCTGAACAA GAGGACCCAG GACTTCTGGG AGGTGCAGCT GGGCATCCCC CACCCCGCTG GCCTGAAGAA GAAGAAGTCT GTGACTGTGC TGGCTGTGGG GGATGCCTAC TTCTCTGTGC CCCTGGATGA GGACTTCAGG AAGTACACTG CCTTCACCAT CCCCTCCATC AACAATGAGA CCCCTGGCAT CAGGTACCAG TACAATGTGC TGCCCCAGGG CTGGAAGGGC TCCCCTGCCA TCTTCCAGTC CTCCATGACC AAGATCCTGG AGCCCTTCAG GAAGCAGAAC 15 CCTGACATTG TGATCTACCA GTACATGGCT GCCCTGTATG TGGGCTCTGA CCTGGAGATT GGGCAGCACA GGACCAAGAT TGAGGAGCTG AGGCAGCACC TGCTGAGGTG GGGCCTGACC ACCCCTGACA AGAAGCACCA GAAGGAGCCC CCCTTCCTGT GGATGGGCTA TGAGCTGCAC CCCGACAAGT GGACTGTGCA GCCCATTGTG CTGCCTGAGA AGGACTCCTG GACTGTGAAT GACATCCAGA AGCTGGTGGG CAAGCTGAAC TGGGCCTCCC AAATCTACCC TGGCATCAAG 20 GTGAGGCAGC TGTGCAAGCT GCTGAGGGGC ACCAAGGCCC TGACTGAGGT GATCCCCCTG ACTGAGGAGG CTGAGCTGGA GCTGGCTGAG AACAGGGAGA TCCTGAAGGA GCCTGTGCAT GGGGTGTACT ATGACCCCTC CAAGGACCTG ATTGCTGAGA TCCAGAAGCA GGGCCAGGGC CAGTGGACCT ACCAAATCTA CCAGGAGCCC TTCAAGAACC TGAAGACTGG CAAGTATGCC AGGATGAGGG GGGCCCACAC CAATGATGTG AAGCAGCTGA CTGAGGCTGT GCAGAAGATC 25 ACCACTGAGT CCATTGTGAT CTGGGGCAAG ACCCCCAAGT TCAAGCTGCC CATCCAGAAG GAGACCTGGG AGACCTGGTG GACTGAGTAC TGGCAGGCCA CCTGGATCCC TGAGTGGGAG TTTGTGAACA CCCCCCCCT GGTGAAGCTG TGGTACCAGC TGGAGAAGGA GCCCATTGTG GGGGCTGAGA CCTTCTATGT GGCTGGGGCT GCCAACAGGG AGACCAAGCT GGGCAAGGCT GGCTATGTGA CCAACAGGGG CAGGCAGAAG GTGGTGACCC TGACTGACAC CACCAACCAG 30 AAGACTGCCC TCCAGGCCAT CTACCTGGCC CTCCAGGACT CTGGCCTGGA GGTGAACATT GTGACTGCCT CCCAGTATGC CCTGGGCATC ATCCAGGCCC AGCCTGATCA GTCTGAGTCT GTGCCTGCCC ACAAGGGCAT TGGGGGCAAT GAGCAGGTGG ACAAGCTGGT GTCTGCTGGC ATCAGGAAGG TGCTGTTCCT GGATGGCATT GACAAGGCCC AGGATGAGCA TGAGAAGTAC 35 CACTCCAACT GGAGGGCTAT GGCCTCTGAC TTCAACCTGC CCCCTGTGGT GGCTAAGGAG

ATTGTGGCCT CCTGTGACAA GTGCCAGCTG AAGGGGAGG CCATGCATGG GCAGGTGGAC
TGCTCCCCTG GCATCTGGCA GCTGGCCTGC ACCCACCTGG AGGGCAAGGT GATCCTGGTG
GCTGTGCATG TGGCCTCCGG CTACATTGAG GCTGAGGTGA TCCCTGCTGA GACAGGCCAG
GAGACTGCCT ACTTCCTGCT GAAGCTGGCT GGCAGGTGC CTGTGAAGAC CATCCACACT
GCCAATGGCT CCAACTTCAC TGGGGCCACA GTGAGGGCTG CCTGCTGGTG GGCTGGCATC
AAGCAGGAGT TTGGCATCCC CTACAACCCC CAGTCCCAGG GGGTGGTGC CTCCATGAAC
AAGGAGCTGA AGAAGATCAT TGGGCAGGTG AGGGACCAGG CTGAGCACCT GAAGACAGCT
GTGCAGATGG CTGTGTTCAT CCACAACTTC AAGAGGAAGG GGGGCATCGG GGGCTACTCC
GCTGGGGAGA GGATTGTGGA CATCATTGCC ACAGACATCC AGACCAAGGA GCTCCAGAAG
CAGATCACCA AGATCCAGAA CTTCAGGGTG TACTACAGGG ACTCCAGGAA CCCCCTGTGG
AAGGGCCCTG CCAAGCTGCT GTGGAAGGGG GAGGGGCCTG TGGTGATCCA GGACAACTCT
GACATCAAGG TGGTGCCCAG GAGGAAGGCC AAGATCATCA GGGACTATGG CAAGCAGATG
GCTGGGGATG ACTGTGTGGC CTCCAGGCAG GATGAGGACT AAAGCCCGGG CAGATCT (SEQ ID
NO:3).

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In order to produce the IA-pol-based adenoviral vaccines of the present 15 invention, inactivation of the enzymatic functions was achieved by replacing a total of nine active site residues from the enzyme subunits with alanine side-chains. As shown in Table 1, all residues that comprise the catalytic triad of the polymerase, namely Asp112, Asp187, and Asp188, were substituted with alanine (Ala) residues (Larder, et al., Nature 1987, 327: 716-717; Larder, et al., 1989, Proc. Natl. Acad. Sci. 20 1989, 86: 4803-4807). Three additional mutations were introduced at Asp445, Glu480 and Asp500 to abolish RNase H activity (Asp551 was left unchanged in this IA Pol construct), with each residue being substituted for an Ala residue, respectively (Davies, et al., 1991, Science 252:, 88-95; Schatz, et al., 1989, FEBS Lett. 257: 311-314; Mizrahi, et al., 1990, Nucl. Acids. Res. 18: pp. 5359-5353). HIV pol integrase 25 function was abolished through three mutations at Asp626, Asp678 and Glu714. Again, each of these residues has been substituted with an Ala residue (Wiskerchen, et al., 1995, J. Virol. 69: 376-386; Leavitt, et al., 1993, J. Biol. Chem. 268: 2113-2119). Amino acid residue Pro3 of SEQ ID NO:4 marks the start of the RT gene. The complete amino acid sequence of IA-Pol is disclosed herein as SEQ ID NO:4 and 30 Figure 17A-C, as follows:

Met Ala Pro Ile Ser Pro Ile Glu Thr Val Pro Val Lys Leu Lys Pro Gly Met Asp Gly Pro Lys Val Lys Gln Trp Pro Leu Thr Glu Glu Lys Ile Lys Ala Leu Val Glu Ile Cys Thr Glu Met Glu Lys Glu Gly Lys Ile Ser Lys Ile Gly Pro Glu Asn Pro Tyr Asn Thr Pro Val Phe Ala Ile Lys Lys Lys Asp Ser Thr Lys Trp Arg Lys Leu Val Asp Phe Arg

Glu Leu Asn Lys Arg Thr Gln Asp Phe Trp Glu Val Gln Leu Gly Ile Pro His Pro Ala Gly Leu Lys Lys Lys Ser Val Thr Val Leu Ala Val Gly Asp Ala Tyr Phe Ser Val Pro Leu Asp Glu Asp Phe Arg Lys Tyr Thr Ala Phe Thr Ile Pro Ser Ile Asn Asn Glu Thr Pro Gly Ile Arg Tyr Gln Tyr Asn Val Leu Pro Gln Gly Trp Lys Gly Ser Pro Ala Ile Phe Gln Ser Ser Met Thr Lys Ile Leu Glu Pro Phe Arg Lys Gln Asn Pro Asp Ile Val Ile Tyr Gln Tyr Met Ala Ala Leu Tyr Val Gly Ser Asp Leu Glu Ile Gly Gln His Arg Thr Lys Ile Glu Glu Leu Arg Gln His Leu Leu Arg Trp Gly Leu Thr Thr Pro Asp Lys Lys His Gln Lys Glu Pro Pro Phe Leu Trp Met Gly Tyr Glu Leu His Pro Asp Lys 10 Trp Thr Val Gln Pro Ile Val Leu Pro Glu Lys Asp Ser Trp Thr Val Asn Asp Ile Gln Lys Leu Val Gly Lys Leu Asn Trp Ala Ser Gln Ile Tyr Pro Gly Ile Lys Val Arg Gln Leu Cys Lys Leu Leu Arg Gly Thr Lys Ala Leu Thr Glu Val Ile Pro Leu Thr Glu Glu Ala Glu Leu Glu Leu Ala Glu Asn Arg Glu Ile Leu Lys Glu Pro Val His Gly Val Tyr 15 Tyr Asp Pro Ser Lys Asp Leu Ile Ala Glu Ile Gln Lys Gln Gly Gln Gly Gln Trp Thr Tyr Gln Ile Tyr Gln Glu Pro Phe Lys Asn Leu Lys Thr Gly Lys Tyr Ala Arg Met Arg Gly Ala His Thr Asn Asp Val Lys Gln Leu Thr Glu Ala Val Gln Lys Ile Thr Thr Glu Ser Ile Val Ile Trp Gly Lys Thr Pro Lys Phe Lys Leu Pro Ile Gln Lys Glu Thr Trp 20 Glu Thr Trp Trp Thr Glu Tyr Trp Gln Ala Thr Trp Ile Pro Glu Trp Glu Phe Val Asn Thr Pro Pro Leu Val Lys Leu Trp Tyr Gln Leu Glu Lys Glu Pro Ile Val Gly Ala Glu Thr Phe Tyr Val Ala Gly Ala Ala Asn Arg Glu Thr Lys Leu Gly Lys Ala Gly Tyr Val Thr Asn Arg Gly Arg Gln Lys Val Val Thr Leu Thr Asp Thr Thr Asn Gln Lys Thr Ala 25 Leu Gln Ala Ile Tyr Leu Ala Leu Gln Asp Ser Gly Leu Glu Val Asn Ile Val Thr Ala Ser Gln Tyr Ala Leu Gly Ile Ile Gln Ala Gln Pro Asp Gln Ser Glu Ser Glu Leu Val Asn Gln Ile Ile Glu Gln Leu Ile Lys Lys Glu Lys Val Tyr Leu Ala Trp Val Pro Ala His Lys Gly Ile Gly Gly Asn Glu Gln Val Asp Lys Leu Val Ser Ala Gly Ile Arg Lys 30 . Val Leu Phe Leu Asp Gly Ile Asp Lys Ala Gln Asp Glu His Glu Lys Tyr His Ser Asn Trp Arg Ala Met Ala Ser Asp Phe Asn Leu Pro Pro Val Val Ala Lys Glu Ile Val Ala Ser Cys Asp Lys Cys Gln Leu Lys Gly Glu Ala Met His Gly Gln Val Asp Cys Ser Pro Gly Ile Trp Gln Leu Ala Cys Thr His Leu Glu Gly Lys Val Ile Leu Val Ala Val His 35 Val Ala Ser Gly Tyr Ile Glu Ala Glu Val Ile Pro Ala Glu Thr Gly

Gln Glu Thr Ala Tyr Phe Leu Leu Lys Leu Ala Gly Arg Trp Pro Val Lys Thr Ile His Thr Ala Asn Gly Ser Asn Phe Thr Gly Ala Thr Val Arg Ala Ala Cys Trp Trp Ala Gly Ile Lys Gln Glu Phe Gly Ile Pro Tyr Asn Pro Gln Ser Gln Gly Val Val Ala Ser Met Asn Lys Glu Leu Lys Lys Ile Ile Gly Gln Val Arg Asp Gln Ala Glu His Leu Lys Thr Ala Val Gln Met Ala Val Phe Ile His Asn Phe Lys Arg Lys Gly Gly Ile Gly Gly Tyr Ser Ala Gly Glu Arg Ile Val Asp Ile Ile Ala Thr Asp Ile Gln Thr Lys Glu Leu Gln Lys Gln Ile Thr Lys Ile Gln Asn Phe Arg Val Tyr Tyr Arg Asp Ser Arg Asn Pro Leu Trp Lys Gly Pro Ala Lys Leu Leu Trp Lys Gly Glu Gly Ala Val Val Ile Gln Asp Asn Ser Asp Ile Lys Val Val Pro Arg Arg Lys Ala Lys Ile Ile Arg Asp Tyr Gly Lys Gln Met Ala Gly Asp Asp Cys Val Ala Ser Arg Gln Asp Glu Asp (SEQ ID NO:4).

As noted above, it will be understood that any combination of the mutations disclosed above may be suitable and therefore be utilized as an IA-pol-based adenoviral HIV vaccine of the present invention, either when administered alone or in a combined modality regime and/or a prime-boost regimen. For example, it may be possible to mutate only 2 of the 3 residues within the respective reverse transcriptase, RNase-H, and integrase coding regions while still abolishing these enzymatic activities. However, the IA-pol construct described above and disclosed as SEQ ID NO:3, as well as the expressed protein (SEQ ID NO:4;) is preferred. It is also preferred that at least one mutation be present in each of the three catalytic domains.

Another aspect of this portion of the invention are codon optimized HIV-1 Pol-based vaccine constructions which comprise a eukaryotic trafficking signal peptide such as from tPA (tissue-type plasminogen activator) or by a leader peptide such as is found in highly expressed mammalian proteins such as immunoglobulin leader peptides. Any functional leader peptide may be tested for efficacy. However, a preferred embodiment of the present invention, as with HIV-1 Nef constructs shown herein, is to provide for a HIV-1 Pol mutant adenoviral vaccine construction wherein the pol coding region or a portion thereof is operatively linked to a leader peptide, preferably a leader peptide from human tPA. In other words, a codon optimized HIV-1 Pol mutant such as IA-Pol (SEQ ID NO:4) may also comprise a leader peptide at the amino terminal portion of the protein, which may effect cellular trafficking and hence, immunogenicity of the expressed protein within the host cell. As noted in Figure 16A-B, a DNA vector which may be utilized to practice the present invention may be modified by known recombinant DNA methodology to contain a leader signal

peptide of interest, such that downstream cloning of the modified HIV-1 protein of interest results in a nucleotide sequence which encodes a modified HIV-1 tPA/Pol protein. In the alternative, as noted above, insertion of a nucleotide sequence which encodes a leader peptide may be inserted into a DNA vector housing the open reading frame for the Pol protein of interest. Regardless of the cloning strategy, the end result is a polynucleotide vaccine which comprises vector components for effective gene expression in conjunction with nucleotide sequences which encode a modified HIV-1 Pol protein of interest, including but not limited to a HIV-1 Pol protein which contains a leader peptide. The amino acid sequence of the human tPA leader utilized herein is as follows: MDAMKRGLCCVLLLCGAVFVSPSEISS (SEQ ID NO:17). Therefore, another aspect of the present invention is to generate HIV-1 Pol-based vaccine constructions which comprise a eukaryotic trafficking signal peptide such as from tPA. To this end, the present invention relates to a DNA molecule which encodes a codon optimized wt-pol DNA construct wherein the protease (PR) activity is deleted and a human tPA leader sequence is fused to the 5' end of the coding region. A DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:5, the open reading frame disclosed herein as SEQ ID NO:6.

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To this end, the present invention relates to a DNA molecule which encodes a codon optimized wt-pol DNA construct wherein the protease (PR) activity is deleted and a human tPA leader sequence is fused to the 5' end of the coding region (herein, "tPA-wt-pol"). A DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:5, the open reading frame being contained from an initiating Met residue at nucleotides 8-10 to a termination codon from nucleotides 2633-2635. SEQ ID NO:5 is as follows:

GATCACCATG GATGCAATGA AGAGAGGGCT CTGCTGTGTG CTGCTGCTGT GTGGAGCAGT
CTTCGTTTCG CCCAGCGAGA TCTCCGCCCC CATCTCCCCC ATTGAGACTG TGCCTGTGAA
GCTGAAGCCT GGCATGGATG GCCCCAAGGT GAAGCAGTGG CCCCTGACTG AGGAGAAGAT
CAAGGCCCTG GTGGAAATCT GCACTGAGAT GGAGAAGGAG GGCAAAATCT CCAAGATTGG
CCCCGAGAAC CCCTACAACA CCCCTGTGTT TGCCATCAAG AAGAAGGACT CCACCAAGTG
GAGGAAGCTG GTGGACTTCA GGGAGCTGAA CAAGAGGACC CAGGACTTCT GGGAGGTGCA
GCTGGGCATC CCCCACCCCG CTGGCCTGAA GAAGAAGAAG TCTGTGACTG TGCTGGATGT
GGGGGATGCC TACTTCTCTG TGCCCCTGGA TGAGGACTTC AGGAAGTACA CTGCCTTCAC
CATCCCCTCC ATCAACAATG AGACCCCTGG CATCAGGTAC CAGTACAATG TGCTGCCCCA
GGGCTGGAAG GGCTCCCCTG CCATCTTCCA GTCCTCCATG ACCAAGATCC TGGAGCCCTT
CAGGAAGCAG AACCCTGACA TTGTGATCTA CCAGTACATG GATGACCTGT ATGTGGGCTC
TGACCTGGAG ATTGGGCAGC ACAGGACCAA GATTGAGGAG CTGAGGCAGC ACCTGCTGAG

GTGGGGCCTG ACCACCCCTG ACAAGAAGCA CCAGAAGGAG CCCCCCTTCC TGTGGATGGG CTATGAGCTG CACCCGACA AGTGGACTGT GCAGCCCATT GTGCTGCCTG AGAAGGACTC CTGGACTGTG AATGACATCC AGAAGCTGGT GGGCAAGCTG AACTGGGCCT CCCAAATCTA CCCTGGCATC AAGGTGAGGC AGCTGTGCAA GCTGCTGAGG GGCACCAAGG CCCTGACTGA GGTGATCCCC CTGACTGAGG AGGCTGAGCT GGAGCTGGCT GAGAACAGGG AGATCCTGAA GGAGCCTGTG CATGGGGTGT ACTATGACCC CTCCAAGGAC CTGATTGCTG AGATCCAGAA GCAGGGCCAG GGCCAGTGGA CCTACCAAAT CTACCAGGAG CCCTTCAAGA ACCTGAAGAC TGGCAAGTAT GCCAGGATGA GGGGGGCCCA CACCAATGAT GTGAAGCAGC TGACTGAGGC TGTGCAGAAG ATCACCACTG AGTCCATTGT GATCTGGGGC AAGACCCCCA AGTTCAAGCT GCCCATCCAG AAGGAGACCT GGGAGACCTG GTGGACTGAG TACTGGCAGG CCACCTGGAT 10 CCCTGAGTGG GAGTTTGTGA ACACCCCCC CCTGGTGAAG CTGTGGTACC AGCTGGAGAA GGAGCCCATT GTGGGGGCTG AGACCTTCTA TGTGGATGGG GCTGCCAACA GGGAGACCAA GCTGGGCAAG GCTGGCTATG TGACCAACAG GGGCAGGCAG AAGGTGGTGA CCCTGACTGA CACCACCAAC CAGAAGACTG AGCTCCAGGC CATCTACCTG GCCCTCCAGG ACTCTGGCCT GGAGGTGAAC ATTGTGACTG ACTCCCAGTA TGCCCTGGGC ATCATCCAGG CCCAGCCTGA 15 TCAGTCTGAG TCTGAGCTGG TGAACCAGAT CATTGAGCAG CTGATCAAGA AGGAGAAGGT GTACCTGGCC TGGGTGCCTG CCCACAAGGG CATTGGGGGC AATGAGCAGG TGGACAAGCT GGTGTCTGCT GGCATCAGGA AGGTGCTGTT CCTGGATGGC ATTGACAAGG CCCAGGATGA GCATGAGAAG TACCACTCCA ACTGGAGGGC TATGGCCTCT GACTTCAACC TGCCCCCTGT GGTGGCTAAG GAGATTGTGG CCTCCTGTGA CAAGTGCCAG CTGAAGGGGG AGGCCATGCA 20 TGGGCAGGTG GACTGCTCCC CTGGCATCTG GCAGCTGGAC TGCACCCACC TGGAGGGCAA GGTGATCCTG GTGGCTGTGC ATGTGGCCTC CGGCTACATT GAGGCTGAGG TGATCCCTGC TGAGACAGGC CAGGAGACTG CCTACTTCCT GCTGAAGCTG GCTGGCAGGT GGCCTGTGAA GACCATCCAC ACTGACAATG GCTCCAACTT CACTGGGGCC ACAGTGAGGG CTGCCTGCTG GTGGGCTGGC ATCAAGCAGG AGTTTGGCAT CCCCTACAAC CCCCAGTCCC AGGGGGTGGT 25 GGAGTCCATG AACAAGGAGC TGAAGAAGAT CATTGGGCAG GTGAGGGACC AGGCTGAGCA CCTGAAGACA GCTGTGCAGA TGGCTGTGTT CATCCACAAC TTCAAGAGGA AGGGGGGCAT CGGGGGCTAC TCCGCTGGGG AGAGGATTGT GGACATCATT GCCACAGACA TCCAGACCAA GGAGCTCCAG AAGCAGATCA CCAAGATCCA GAACTTCAGG GTGTACTACA GGGACTCCAG 30 GAACCCCTG TGGAAGGCC CTGCCAAGCT GCTGTGGAAG GGGGAGGGGG CTGTGGTGAT CCAGGACAAC TCTGACATCA AGGTGGTGCC CAGGAGGAAG GCCAAGATCA TCAGGGACTA TGGCAAGCAG ATGGCTGGGG ATGACTGTGT GGCCTCCAGG CAGGATGAGG ACTAAAGCCC GGGCAGATCT (SEQ ID NO:5).

The open reading frame of the wild type tPA-pol construct disclosed as SEQ ID NO:5 contains 875 amino acids, disclosed herein as SEQ ID NO:6, as follows:

Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Cys Gly

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Ala Val Phe Val Ser Pro Ser Glu Ile Ser Ala Pro Ile Ser Pro Ile Glu Thr Val Pro Val Lys Leu Lys Pro Gly Met Asp Gly Pro Lys Val Lys Gln Trp Pro Leu Thr Glu Glu Lys Ile Lys Ala Leu Val Glu Ile Cys Thr Glu Met Glu Lys Glu Gly Lys Ile Ser Lys Ile Gly Pro Glu Asn Pro Tyr Asn Thr Pro Val Phe Ala Ile Lys Lys Lys Asp Ser Thr 5 Lys Trp Arg Lys Leu Val Asp Phe Arg Glu Leu Asn Lys Arg Thr Gln Asp Phe Trp Glu Val Gln Leu Gly Ile Pro His Pro Ala Gly Leu Lys Lys Lys Ser Val Thr Val Leu Asp Val Gly Asp Ala Tyr Phe Ser Val Pro Leu Asp Glu Asp Phe Arg Lys Tyr Thr Ala Phe Thr Ile Pro Ser Ile Asn Asn Glu Thr Pro Gly Ile Arg Tyr Gln Tyr Asn Val Leu 10 Pro Gln Gly Trp Lys Gly Ser Pro Ala Ile Phe Gln Ser Ser Met Thr Lys Ile Leu Glu Pro Phe Arg Lys Gln Asn Pro Asp Ile Val Ile Tyr Gln Tyr Met Asp Asp Leu Tyr Val Gly Ser Asp Leu Glu Ile Gly Gln His Arg Thr Lys Ile Glu Glu Leu Arg Gln His Leu Leu Arg Trp Gly Leu Thr Thr Pro Asp Lys Lys His Gln Lys Glu Pro Pro Phe Leu Trp 15 Met Gly Tyr Glu Leu His Pro Asp Lys Trp Thr Val Gln Pro Ile Val Leu Pro Glu Lys Asp Ser Trp Thr Val Asn Asp Ile Gln Lys Leu Val Gly Lys Leu Asn Trp Ala Ser Gln Ile Tyr Pro Gly Ile Lys Val Arg Gln Leu Cys Lys Leu Leu Arg Gly Thr Lys Ala Leu Thr Glu Val Ile Pro Leu Thr Glu Glu Ala Glu Leu Glu Leu Ala Glu Asn Arg Glu Ile 20 Leu Lys Glu Pro Val His Gly Val Tyr Tyr Asp Pro Ser Lys Asp Leu Ile Ala Glu Ile Gln Lys Gln Gly Gln Gly Gln Trp Thr Tyr Gln Ile Tyr Gln Glu Pro Phe Lys Asn Leu Lys Thr Gly Lys Tyr Ala Arg Met Arg Gly Ala His Thr Asn Asp Val Lys Gln Leu Thr Glu Ala Val Gln Lys Ile Thr Thr Glu Ser Ile Val Ile Trp Gly Lys Thr Pro Lys Phe 25 Lys Leu Pro Ile Gln Lys Glu Thr Trp Glu Thr Trp Trp Thr Glu Tyr Trp Gln Ala Thr Trp Ile Pro Glu Trp Glu Phe Val Asn Thr Pro Pro Leu Val Lys Leu Trp Tyr Gln Leu Glu Lys Glu Pro Ile Val Gly Ala Glu Thr Phe Tyr Val Asp Gly Ala Ala Asn Arg Glu Thr Lys Leu Gly Lys Ala Gly Tyr Val Thr Asn Arg Gly Arg Gln Lys Val Val Thr Leu 30 Thr Asp Thr Thr Asn Gln Lys Thr.Glu Leu Gln Ala Ile Tyr Leu Ala Leu Gln Asp Ser Gly Leu Glu Val Asn Ile Val Thr Asp Ser Gln Tyr Ala Leu Gly Ile Ile Gln Ala Gln Pro Asp Gln Ser Glu Ser Glu Leu Val Asn Gln Ile Ile Glu Gln Leu Ile Lys Lys Glu Lys Val Tyr Leu Ala Trp Val Pro Ala His Lys Gly Ile Gly Gly Asn Glu Gln Val Asp 35 Lys Leu Val Ser Ala Gly Ile Arg Lys Val Leu Phe Leu Asp Gly Ile

Asp Lys Ala Gln Asp Glu His Glu Lys Tyr His Ser Asn Trp Arg Ala Met Ala Ser Asp Phe Asn Leu Pro Pro Val Val Ala Lys Glu Ile Val Ala Ser Cys Asp Lys Cys Gln Leu Lys Gly Glu Ala Met His Gly Gln Val Asp Cys Ser Pro Gly Ile Trp Gln Leu Asp Cys Thr His Leu Glu Gly Lys Val Ile Leu Val Ala Val His Val Ala Ser Gly Tyr Ile Glu Ala Glu Val Ile Pro Ala Glu Thr Gly Gln Glu Thr Ala Tyr Phe Leu Leu Lys Leu Ala Gly Arg Trp Pro Val Lys Thr Ile His Thr Asp Asn Gly Ser Asn Phe Thr Gly Ala Thr Val Arg Ala Ala Cys Trp Trp Ala Gly Ile Lys Gln Glu Phe Gly Ile Pro Tyr Asn Pro Gln Ser Gln Gly 10 Val Val Glu Ser Met Asn Lys Glu Leu Lys Lys Ile Ile Gly Gln Val Arg Asp Gln Ala Glu His Leu Lys Thr Ala Val Gln Met Ala Val Phe Ile His Asn Phe Lys Arg Lys Gly Gly Ile Gly Gly Tyr Ser Ala Gly Glu Arg Ile Val Asp Ile Ile Ala Thr Asp Ile Gln Thr Lys Glu Leu Gln Lys Gln Ile Thr Lys Ile Gln Asn Phe Arg Val Tyr Tyr Arg Asp 15 Ser Arg Asn Pro Leu Trp Lys Gly Pro Ala Lys Leu Leu Trp Lys Gly Glu Gly Ala Val Val Ile Gln Asp Asn Ser Asp Ile Lys Val Val Pro Arg Arg Lys Ala Lys Ile Ile Arg Asp Tyr Gly Lys Gln Met Ala Gly Asp Asp Cys Val Ala Ser Arg Gln Asp Glu Asp (SEQ ID NO:6).

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The present invention also relates to a codon optimized HIV-1 Pol mutant contained within a recombinant adenoviral vector such as IA-Pol (SEQ ID NO:4) which comprises a leader peptide at the amino terminal portion of the protein, which may effect cellular trafficking and hence, immunogenicity of the expressed protein within the host cell. Any such adenoviral-based HIV-1 DNA pol mutant disclosed in the above paragraphs is suitable for fusion downstream of a leader peptide, such as a leader peptide including but not limited to the human tPA leader sequence. Therefore, any such leader peptide-based HIV-1 pol mutant construct may include but is not limited to a mutated DNA molecule which effectively alters the catalytic activity of the RT, RNase and/or IN region of the expressed protein, resulting in at least substantially decreased enzymatic activity one or more of the RT, RNase H and/or IN functions of HIV-1 Pol. In a preferred embodiment of this portion of the invention, a leader peptide/HIV-1 DNA pol construct contains a mutation or mutations within the Pol coding region which effectively abolishes RT, RNase H and IN activity. An especially preferable HIV-1 DNA pol construct is a DNA molecule which contains at least one point mutation which alters the active site and catalytic activity within the RT, RNase H and IN domains of Pol, such that each activity is at least substantially abolished, and preferably totally abolished. Such a HIV-1 Pol mutant will most likely

comprise at least one point mutation in or around each catalytic domain responsible for RT, RNase H and IN activity, respectfully. An especially preferred embodiment of this portion of the invention relates to a human tPA leader fused to the IA-Pol protein comprising the nine mutations shown in Table 1. The DNA molecule is disclosed 5 herein as SEQ ID NO:7 and the expressed tPA-IA Pol protein comprises a fusion junction as shown in Figure 18. The complete amino acid sequence of the expressed protein is set forth in SEQ ID NO:8. To this end, SEQ ID NO:7 discloses the nucleotide sequence which codes for a human tPA leader fused to the IA Pol protein comprising the nine mutations shown in Table 1 (herein, "tPA-opt-IApol"). The open 10 reading frame begins with the initiating Met (nucleotides 8-10) and terminates with a "TAA" codon at nucleotides 2633-2635. The nucleotide sequence encoding tPA-IAPol is also disclosed as follows: GATCACCATG GATGCAATGA AGAGAGGGCT CTGCTGTGTG CTGCTGCTGT GTGGAGCAGT CTTCGTTTCG CCCAGCGAGA TCTCCGCCCC CATCTCCCCC ATTGAGACTG TGCCTGTGAA 15 GCTGAAGCCT GGCATGGATG GCCCCAAGGT GAAGCAGTGG CCCCTGACTG AGGAGAAGAT CAAGGCCCTG GTGGAAATCT GCACTGAGAT GGAGAAGGAG GGCAAAATCT CCAAGATTGG CCCCGAGAC CCCTACAACA CCCCTGTGTT TGCCATCAAG AAGAAGGACT CCACCAAGTG GAGGAAGCTG GTGGACTTCA GGGAGCTGAA CAAGAGGACC CAGGACTTCT GGGAGGTGCA GCTGGGCATC CCCCACCCG CTGGCCTGAA GAAGAAGAAG TCTGTGACTG TGCTGGCTGT 20 GGGGGATGCC TACTTCTCTG TGCCCCTGGA TGAGGACTTC AGGAAGTACA CTGCCTTCAC CATCCCCTCC ATCAACAATG AGACCCCTGG CATCAGGTAC CAGTACAATG TGCTGCCCCA GGGCTGGAAG GGCTCCCCTG CCATCTTCCA GTCCTCCATG ACCAAGATCC TGGAGCCCTT CAGGAAGCAG AACCCTGACA TTGTGATCTA CCAGTACATG GCTGCCCTGT ATGTGGGCTC TGACCTGGAG ATTGGGCAGC ACAGGACCAA GATTGAGGAG CTGAGGCAGC ACCTGCTGAG 25 GTGGGGCCTG ACCACCCCTG ACAAGAAGCA CCAGAAGGAG CCCCCCTTCC TGTGGATGGG CTATGAGCTG CACCCCGACA AGTGGACTGT GCAGCCCATT GTGCTGCCTG AGAAGGACTC CTGGACTGTG AATGACATCC AGAAGCTGGT GGGCAAGCTG AACTGGGCCT CCCAAATCTA CCCTGGCATC AAGGTGAGGC AGCTGTGCAA GCTGCTGAGG GGCACCAAGG CCCTGACTGA GGTGATCCCC CTGACTGAGG AGGCTGAGCT GGAGCTGGCT GAGAACAGGG AGATCCTGAA 30 GGAGCCTGTG CATGGGGTGT ACTATGACCC CTCCAAGGAC CTGATTGCTG AGATCCAGAA GCAGGGCCAG GGCCAGTGGA CCTACCAAAT CTACCAGGAG CCCTTCAAGA ACCTGAAGAC TGGCAAGTAT GCCAGGATGA GGGGGGCCCA CACCAATGAT GTGAAGCAGC TGACTGAGGC TGTGCAGAAG ATCACCACTG AGTCCATTGT GATCTGGGGC AAGACCCCCA AGTTCAAGCT GCCCATCCAG AAGGAGACCT GGGAGACCTG GTGGACTGAG TACTGGCAGG CCACCTGGAT 35 CCCTGAGTGG GAGTTTGTGA ACACCCCCC CCTGGTGAAG CTGTGGTACC AGCTGGAGAA GGAGCCCATT GTGGGGGCTG AGACCTTCTA TGTGGCTGGG GCTGCCAACA GGGAGACCAA

GCTGGGCAAG GCTGGCTATG TGACCAACAG GGGCAGGCAG AAGGTGGTGA CCCTGACTGA CACCACCAAC CAGAAGACTG CCCTCCAGGC CATCTACCTG GCCCTCCAGG ACTCTGGCCT GGAGGTGAAC ATTGTGACTG CCTCCCAGTA TGCCCTGGGC ATCATCCAGG CCCAGCCTGA TCAGTCTGAG TCTGAGCTGG TGAACCAGAT CATTGAGCAG CTGATCAAGA AGGAGAAGGT GTACCTGGCC TGGGTGCCTG CCCACAAGGG CATTGGGGGC AATGAGCAGG TGGACAAGCT GGTGTCTGCT GGCATCAGGA AGGTGCTGTT CCTGGATGGC ATTGACAAGG CCCAGGATGA GCATGAGAAG TACCACTCCA ACTGGAGGGC TATGGCCTCT GACTTCAACC TGCCCCCTGT GGTGGCTAAG GAGATTGTGG CCTCCTGTGA CAAGTGCCAG CTGAAGGGGG AGGCCATGCA TGGGCAGGTG GACTGCTCCC CTGGCATCTG GCAGCTGGCC TGCACCCACC TGGAGGGCAA GGTGATCCTG GTGGCTGTGC ATGTGGCCTC CGGCTACATT GAGGCTGAGG TGATCCCTGC 10 TGAGACAGGC CAGGAGACTG CCTACTTCCT GCTGAAGCTG GCTGGCAGGT GGCCTGTGAA GACCATCCAC ACTGCCAATG GCTCCAACTT CACTGGGGCC ACAGTGAGGG CTGCCTGCTG GTGGGCTGGC ATCAAGCAGG AGTTTGGCAT CCCCTACAAC CCCCAGTCCC AGGGGGTGGT GGCCTCCATG AACAAGGAGC TGAAGAAGAT CATTGGGCAG GTGAGGGACC AGGCTGAGCA CCTGAAGACA GCTGTGCAGA TGGCTGTGTT CATCCACAAC TTCAAGAGGA AGGGGGGCAT 15 CGGGGGCTAC TCCGCTGGGG AGAGGATTGT GGACATCATT GCCACAGACA TCCAGACCAA GGAGCTCCAG AAGCAGATCA CCAAGATCCA GAACTTCAGG GTGTACTACA GGGACTCCAG GAACCCCCTG TGGAAGGGCC CTGCCAAGCT GCTGTGGAAG GGGGAGGGGG CTGTGGTGAT CCAGGACAAC TCTGACATCA AGGTGGTGCC CAGGAGGAAG GCCAAGATCA TCAGGGACTA TGGCAAGCAG ATGGCTGGGG ATGACTGTGT GGCCTCCAGG CAGGATGAGG ACTAAAGCCC 20 GGGCAGATCT (SEQ ID NO:7).

The open reading frame of the tPA-IA-pol construct disclosed as SEQ ID NO:7 contains 875 amino acids, disclosed herein as tPA-IA-Pol and SEQ ID NO:8, as follows:

Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Leu Cys Gly
Ala Val Phe Val Ser Pro Ser Glu Ile Ser Ala Pro Ile Ser Pro Ile
Glu Thr Val Pro Val Lys Leu Lys Pro Gly Met Asp Gly Pro Lys Val
Lys Gln Trp Pro Leu Thr Glu Glu Lys Ile Lys Ala Leu Val Glu Ile
Cys Thr Glu Met Glu Lys Glu Gly Lys Ile Ser Lys Ile Gly Pro Glu

Asn Pro Tyr Asn Thr Pro Val Phe Ala Ile Lys Lys Lys Asp Ser Thr
Lys Trp Arg Lys Leu Val Asp Phe Arg Glu Leu Asn Lys Arg Thr Gln
Asp Phe Trp Glu Val Gln Leu Gly Ile Pro His Pro Ala Gly Leu Lys
Lys Lys Lys Ser Val Thr Val Leu Ala Val Gly Asp Ala Tyr Phe Ser
Val Pro Leu Asp Glu Asp Phe Arg Lys Tyr Thr Ala Phe Thr Ile Pro
Ser Ile Asn Asn Glu Thr Pro Gly Ile Arg Tyr Gln Tyr Asn Val Leu
Pro Gln Gly Trp Lys Gly Ser Pro Ala Ile Phe Gln Ser Ser Met Thr

Lys Ile Leu Glu Pro Phe Arg Lys Gln Asn Pro Asp Ile Val Ile Tyr Gln Tyr Met Ala Ala Leu Tyr Val Gly Ser Asp Leu Glu Ile Gly Gln His Arg Thr Lys Ile Glu Glu Leu Arg Gln His Leu Leu Arg Trp Gly Leu Thr Thr Pro Asp Lys Lys His Gln Lys Glu Pro Pro Phe Leu Trp Met Gly Tyr Glu Leu His Pro Asp Lys Trp Thr Val Gln Pro Ile Val Leu Pro Glu Lys Asp Ser Trp Thr Val Asn Asp Ile Gln Lys Leu Val Gly Lys Leu Asn Trp Ala Ser Gln Ile Tyr Pro Gly Ile Lys Val Arg Gln Leu Cys Lys Leu Leu Arg Gly Thr Lys Ala Leu Thr Glu Val Ile Pro Leu Thr Glu Glu Ala Glu Leu Glu Leu Ala Glu Asn Arg Glu Ile Leu Lys Glu Pro Val His Gly Val Tyr Tyr Asp Pro Ser Lys Asp Leu 10 Ile Ala Glu Ile Gln Lys Gln Gly Gln Gly Gln Trp Thr Tyr Gln Ile Tyr Gln Glu Pro Phe Lys Asn Leu Lys Thr Gly Lys Tyr Ala Arg Met Arg Gly Ala His Thr Asn Asp Val Lys Gln Leu Thr Glu Ala Val Gln Lys Ile Thr Thr Glu Ser Ile Val Ile Trp Gly Lys Thr Pro Lys Phe Lys Leu Pro Ile Gln Lys Glu Thr Trp Glu Thr Trp Trp Thr Glu Tyr 15 Trp Gln Ala Thr Trp Ile Pro Glu Trp Glu Phe Val Asn Thr Pro Pro Leu Val Lys Leu Trp Tyr Gln Leu Glu Lys Glu Pro Ile Val Gly Ala Glu Thr Phe Tyr Val Ala Gly Ala Ala Asn Arg Glu Thr Lys Leu Gly Lys Ala Gly Tyr Val Thr Asn Arg Gly Arg Gln Lys Val Val Thr Leu Thr Asp Thr Thr Asn Gln Lys Thr Ala Leu Gln Ala Ile Tyr Leu Ala 20 Leu Gln Asp Ser Gly Leu Glu Val Asn Ile Val Thr Ala Ser Gln Tyr Ala Leu Gly Ile Ile Gln Ala Gln Pro Asp Gln Ser Glu Ser Glu Leu Val Asn Gln Ile Ile Glu Gln Leu Ile Lys Lys Glu Lys Val Tyr Leu Ala Trp Val Pro Ala His Lys Gly Ile Gly Gly Asn Glu Gln Val Asp Lys Leu Val Ser Ala Gly Ile Arg Lys Val Leu Phe Leu Asp Gly Ile 25 Asp Lys Ala Gln Asp Glu His Glu Lys Tyr His Ser Asn Trp Arg Ala Met Ala Ser Asp Phe Asn Leu Pro Pro Val Val Ala Lys Glu Ile Val Ala Ser Cys Asp Lys Cys Gln Leu Lys Gly Glu Ala Met His Gly Gln Val Asp Cys Ser Pro Gly Ile Trp Gln Leu Ala Cys Thr His Leu Glu 30 Gly Lys Val Ile Leu Val Ala Val His Val Ala Ser Gly Tyr Ile Glu Ala Glu Val Ile Pro Ala Glu Thr Gly Gln Glu Thr Ala Tyr Phe. Leu Leu Lys Leu Ala Gly Arg Trp Pro Val Lys Thr Ile His Thr Ala Asn Gly Ser Asn Phe Thr Gly Ala Thr Val Arg Ala Ala Cys Trp Trp Ala Gly Ile Lys Gln Glu Phe Gly Ile Pro Tyr Asn Pro Gln Ser Gln Gly Val Val Ala Ser Met Asn Lys Glu Leu Lys Lys Ile Ile Gly Gln Val 35 Arg Asp Gln Ala Glu His Leu Lys Thr Ala Val Gln Met Ala Val Phe

Ile His Asn Phe Lys Arg Lys Gly Gly Ile Gly Gly Tyr Ser Ala Gly Glu Arg Ile Val Asp Ile Ile Ala Thr Asp Ile Gln Thr Lys Glu Leu Gln Lys Gln Ile Thr Lys Ile Gln Asn Phe Arg Val Tyr Tyr Arg Asp Ser Arg Asn Pro Leu Trp Lys Gly Pro Ala Lys Leu Leu Trp Lys Gly Glu Gly Ala Val Val Ile Gln Asp Asn Ser Asp Ile Lys Val Val Pro Arg Arg Lys Ala Lys Ile Ile Arg Asp Tyr Gly Lys Gln Met Ala Gly Asp Asp Cys Val Ala Ser Arg Gln Asp Glu Asp (SEQ ID NO:8).

EXAMPLE 18

CODON OPTIMIZED HIV-1 NEF AND CODON OPTIMIZED HIV-1 NEF MODIFICATIONS

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Codon optimized version of HIV-1 Nef and HIV-1 Nef modifications are essentially as described in U.S. Application Serial No. 09/738,782, filed December 15, 2000 and PCT International Application PCT/US00/34162, also filed December 15, 2000, both documents which are hereby incorporated by reference. As 15 disclosed within the above-mentioned documents, particular embodiments of codon optimized Nef and Nef modifications relate to a DNA molecule encoding HIV-1 Nef from the HIV-1 jfrl isolate wherein the codons are optimized for expression in a mammalian system such as a human. The DNA molecule which encodes this protein 20 is disclosed herein as SEQ ID NO:9, while the expressed open reading frame is disclosed herein as SEQ ID NO:10. Another embodiment of Nef-based coding regions for use in the adenoviral vectors of the present invention comprise a codon optimized DNA molecule encoding a protein containing the human plasminogen activator (tpa) leader peptide fused with the NH2-terminus of the HIV-1 Nef polypeptide. The DNA molecule which encodes this protein is disclosed herein as 25 SEQ ID NO:11, while the expressed open reading frame is disclosed herein as SEQ ID NO:12. Another modified Nef optimized coding region relates to a DNA molecule encoding optimized HIV-1 Nef wherein the open reading frame codes for modifications at the amino terminal myristylation site (Gly-2 to Ala-2) and 30 substitution of the Leu-174-Leu-175 dileucine motif to Ala-174-Ala-175, herein described as opt nef (G2A, LLAA). The DNA molecule which encodes this protein is disclosed herein as SEQ ID NO:13, while the expressed open reading frame is disclosed herein as SEQ ID NO:14. An additional embodiment relates to a DNA molecule encoding optimized HIV-1 Nef wherein the amino terminal myristylation site and dileucine motif have been deleted, as well as comprising a tPA leader peptide. 35 This DNA molecule, opt tpanef (LLAA), comprises an open reading frame which

encodes a Nef protein containing a tPA leader sequence fused to amino acid residue 6-216 of HIV-1 Nef (ifrl), wherein Leu-174 and Leu-175 are substituted with Ala-174 and Ala-175, herein referred to as opt tpanef (LLAA) is disclosed herein as SEQ ID NO:15, while the expressed open reading frame is disclosed herein as SEQ ID NO:16.

As disclosed in the above-identified documents (U.S. Application Serial No. 09/738,782 and PCT International Application PCT/US00/34162) and reiterated herein, the following nef-based nucleotide and amino acid sequences which comprise the respective open reading frame are as follows:

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The nucleotide sequence of the codon optimized version of HIV-1 jrfl nef gene is disclosed herein as SEQ ID NO:9, as shown herein: GATCTGCCAC CATGGGCGGC AAGTGGTCCA AGAGGTCCGT GCCCGGCTGG TCCACCGTGA GGGAGAGGAT GAGGAGGGCC GAGCCCGCCG CCGACAGGGT GAGGAGGACC GAGCCCGCCG CCGTGGGCGT GGGCGCCGTG TCCAGGGACC TGGAGAAGCA CGGCGCCATC ACCTCCTCCA ACACCGCCGC CACCAACGCC GACTGCGCCT GGCTGGAGGC CCAGGAGGAC GAGGAGGTGG GCTTCCCCGT GAGGCCCCAG GTGCCCCTGA GGCCCATGAC CTACAAGGGC GCCGTGGACC TGTCCCACTT CCTGAAGGAG AAGGGCGGCC TGGAGGGCCT GATCCACTCC CAGAAGAGGC AGGACATCCT GGACCTGTGG GTGTACCACA CCCAGGGCTA CTTCCCCGAC TGGCAGAACT ACACCCCGG CCCCGGCATC AGGTTCCCCC TGACCTTCGG CTGGTGCTTC AAGCTGGTGC CCGTGGAGCC CGAGAAGGTG GAGGAGGCCA ACGAGGGCGA GAACAACTGC CTGCTGCACC CCATGTCCCA GCACGGCATC GAGGACCCCG AGAAGGAGGT GCTGGAGTGG AGGTTCGACT CCAAGCTGGC CTTCCACCAC GTGGCCAGGG AGCTGCACCC CGAGTACTAC AAGGACTGCT AAAGCCCGGG C (SEQ ID NO:9).

Preferred codon usage is as follows: Met (ATG), Gly (GGC), Lys (AAG), Trp (TGG), Ser (TCC), Arg (AGG), Val (GTG), Pro (CCC), Thr (ACC), Glu (GAG); Leu (CTG), His (CAC), Ile (ATC), Asn (AAC), Cys (TGC), Ala (GCC), Gln (CAG), Phe (TTC) and Tyr (TAC). For an additional discussion relating to mammalian (human) codon optimization, see WO 97/31115 (PCT/US97/02294), which is hereby incorporated by reference. See also Figure 19A-B for a comparion of wild type vs. codon optimized nucleotides comprising the open reading frame of HIV-Nef.

The open reading frame for SEQ ID NO:9 above comprises an initiating methionine residue at nucleotides 12-14 and a "TAA" stop codon from nucleotides 660-662. The open reading frame of SEQ ID NO:9 provides for a 216 amino acid HTV-1 Nef protein expressed through utilization of a codon optimized DNA vaccine vector. The 216 amino acid HIV-1 Nef (jfrl) protein is disclosed herein as SEQ ID 35 NO:10, and as follows:

Met Gly Gly Lys Trp Ser Lys Arg Ser Val Pro Gly Trp Ser Thr Val

Arg Glu Arg Met Arg Arg Ala Glu Pro Ala Ala Asp Arg Val Arg Arg Arg Thr Glu Pro Ala Ala Val Gly Val Gly Ala Val Ser Arg Asp Leu Glu Lys His Gly Ala Ile Thr Ser Ser Asn Thr Ala Ala Thr Asn Ala Asp Cys Ala Trp Leu Glu Ala Gln Glu Asp Glu Glu Val Gly Phe Pro Val Arg Pro Gln Val Pro Leu Arg Pro Met Thr Tyr Lys Gly Ala Val Asp Leu Ser His Phe Leu Lys Glu Lys Gly Gly Leu Glu Gly Leu Ile His Ser Gln Lys Arg Gln Asp Ile Leu Asp Leu Trp Val Tyr His Thr Gln Gly Tyr Phe Pro Asp Trp Gln Asn Tyr Thr Pro Gly Pro Gly Ile Arg Phe Pro Leu Thr Phe Gly Trp Cys Phe Lys Leu Val Pro Val Glu Pro Glu Lys Val Glu Glu Ala Asn Glu Gly Glu Asn Asn Cys Leu Leu His Pro Met Ser Gln His Gly Ile Glu Asp Pro Glu Lys Glu Val Leu Glu Trp Arg Phe Asp Ser Lys Leu Ala Phe His His Val Ala Arg Glu Leu His Pro Glu Tyr Tyr Lys Asp Cys (SEQ ID NO:10).

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HIV-1 Nef is a 216 amino acid cytosolic protein which associates with the inner surface of the host cell plasma membrane through myristylation of Gly-2 15 (Franchini et al., 1986, Virology 155: 593-599). While not all possible Nef functions have been elucidated, it has become clear that correct trafficking of Nef to the inner plasma membrane promotes viral replication by altering the host intracellular environment to facilitate the early phase of the HIV-1 life cycle and by increasing the infectivity of progeny viral particles. In one aspect of the invention regarding 20 codon-optimized, protein-modified polypeptides, the nef-encoding region of the adenovirus vector of the present invention is modified to contain a nucleotide sequence which encodes a heterologous leader peptide such that the amino terminal region of the expressed protein will contain the leader peptide. The diversity of function that typifies eukaryotic cells depends upon the structural differentiation of 25 their membrane boundaries. To generate and maintain these structures, proteins must be transported from their site of synthesis in the endoplasmic reticulum to predetermined destinations throughout the cell. This requires that the trafficking proteins display sorting signals that are recognized by the molecular machinery responsible for route selection located at the access points to the main trafficking 30 pathways. Sorting decisions for most proteins need to be made only once as they traverse their biosynthetic pathways since their final destination, the cellular location at which they perform their function, becomes their permanent residence. Maintenance of intracellular integrity depends in part on the selective sorting and accurate transport of proteins to their correct destinations. Defined sequence motifs 35 exist in proteins which can act as 'address labels'. A number of sorting signals have

been found associated with the cytoplasmic domains of membrane proteins. An effective induction of CTL responses often required sustained, high level endogenous expression of an antigen. As membrane-association via myristylation is an essential requirement for most of Nef's function, mutants lacking myristylation, by glycine-to-alanine change, change of the dileucine motif and/or by substitution with a tpa leader sequence as described herein, will be functionally defective, and therefore will have improved safety profile compared to wild-type Nef for use as an HIV-1 vaccine component.

In another embodiment of this portion of the invention, either the DNA vector or the HIV-1 nef nucleotide sequence is modified to include the human tissue-specific plasminogen activator (tPA) leader. As shown in Figure 16A-B, a DNA vector may be modified by known recombinant DNA methodology to contain a leader signal peptide of interest, such that downstream cloning of the modified HIV-1 protein of interest results in a nucleotide sequence which encodes a modified HIV-1 tPA/Nef protein. In the alternative, as noted above, insertion of a nucleotide sequence which encodes a leader peptide may be inserted into a DNA vector housing the open reading frame for the Nef protein of interest. Regardless of the cloning strategy, the end result is a polynucleotide vaccine which comprises vector components for effective gene expression in conjunction with nucleotide sequences which encode a modified HIV-1 Nef protein of interest, including but not limited to a HIV-1 Nef protein which contains a leader peptide. The amino acid sequence of the human tPA leader utilized herein is as follows: MDAMKRGLCCVLLLCGAVFVSPSEISS (SEQ ID NO:17).

It has been shown that myristylation of Gly-2 in conjunction with a dileucine motif in the carboxy region of the protein is essential for Nef-induced down regulation of CD4 (Aiken et al., 1994, Cell 76: 853-864) via endocytosis. It has also been shown that Nef expression promotes down regulation of MHCI (Schwartz et al., 1996, Nature Medicine 2(3): 338-342) via endocytosis. The present invention relates in part to DNA vaccines which encode modified Nef proteins altered in trafficking and/or functional properties. The modifications introduced into the adenoviral vector HIV vaccines of the present invention include but are not limited to additions, deletions or substitutions to the nef open reading frame which results in the expression of a modified Nef protein which includes an amino terminal leader peptide, modification or deletion of the amino terminal myristylation site, and modification or deletion of the dileucine motif within the Nef protein and which alter function within the infected host cell. Therefore, a central theme of the DNA molecules and recombinant adenoviral HIV vaccines of the present invention is (1)

host administration and intracellular delivery of a codon optimized nef-based adenoviral HIV vaccine; (2) expression of a modified Nef protein which is immunogenic in terms of eliciting both CTL and Th responses; and, (3) inhibiting or at least altering known early viral functions of Nef which have been shown to promote HIV-1 replication and load within an infected host. Therefore, the nef coding region may be altered, resulting in a DNA vaccine which expresses a modified Nef protein wherein the amino terminal Gly-2 myristylation residue is either deleted or modified to express alternate amino acid residues. Also, the nef coding region may be altered so as to result in a DNA vaccine which expresses a modified Nef protein wherein the dileucine motif is either deleted or modified to express alternate amino acid residues. In addition, the adenoviral vector HIV vaccines of the present invention also relate to an isolated DNA molecule, regardless of codon usage, which expresses a wild type or modified Nef protein as described herein, including but not limited to modified Nef proteins which comprise a deletion or substitution of Gly 2, a deletion or substitution of Leu 174 and Leu 175 and/or inclusion of a leader sequence.

Therefore, specific Nef-based constructs further include the following, as exemplification's and not limitations. For example, the present invention relates to an adenoviral vector vaccine which encodes modified forms of HIV-1, an open reading frame which encodes a Nef protein which comprises a tPA leader sequence fused to amino acid residue 6-216 of HIV-1 Nef (jfrl) is referred to herein as opt tpanef. The nucleotide sequence comprising the open reading frame of opt tpanef is disclosed herein as SEQ ID NO:11, as shown below:

CATGGATGCA ATGAAGAGA GGCTCTGCTG TGTGCTGCTG CTGTGTGGAG CAGTCTTCGT
TTCGCCCAGC GAGATCTCCT CCAAGAGGTC CGTGCCCGGC TGGTCCACCG TGAGGGAGAG
GATGAGGAGG GCCGAGCCCG CCGCCGACAG GGTGAGGAGG ACCGAGCCCG CCGCCGTGGG
CGTGGGCGCC GTGTCCAGGG ACCTGGAGAA GCACGGCGCC ATCACCTCCT CCAACACCGC
CGCCACCAAC GCCGACTGCG CCTGGCTGGA GGCCCAGGAG GACGAGGAGG TGGGCTTCCC
CGTGAGGCCC CAGGTGCCCC TGAGGCCCAT GACCTACAAG GGCGCCGTGG ACCTGTCCCA
CTTCCTGAAG GAGAAGGGCG GCCTGGAGGG CCTGATCCAC TCCCAGAAGA GGCAGGACAT
CCTGGACCTG TGGGTGTACC ACACCCAGGG CTACTTCCCC GACTGGCAGA ACTACACCCC
CGGCCCCGGC ATCAGGTTCC CCCTGACCTT CGGCTGGTGC TTCAAGCTGG TGCCCGTGGA
GCCCGAGAAG GTGGAGGAGC CCAACGAGGG CGAGAACAAC TGCCTGCTGC ACCCCATGTC
CCAGCACGCC ATCGAGGACC CCGAGAAGGA GGTGCTGGAG TGGAGGTTCG ACCCCATGTC
GGCCTTCCAC CACGTGGCCA GGGAGCTGCA CCCCGAGTAC TACAAGGACT GCTAAAGCC

The open reading frame for SEQ ID NO:11 comprises an initiating methionine

residue at nucleotides 2-4 and a "TAA" stop codon from nucleotides 713-715. The open reading frame of SEQ ID NO:3 provides for a 237 amino acid HIV-1 Nef protein which comprises a tPA leader sequence fused to amino acids 6-216 of HIV-1 Nef, including the dileucine motif at amino acid residues 174 and 175. This 237 amino acid tPA/Nef (jfrl) fusion protein is disclosed herein as SEQ ID NO:12, and is shown as follows:

Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Cys Gly Ala Val Phe Val Ser Pro Ser Glu Ile Ser Ser Lys Arg Ser Val Pro Gly Trp Ser Thr Val Arg Glu Arg Met Arg Arg Ala Glu Pro Ala Ala Asp Arg Val Arg Arg Thr Glu Pro Ala Ala Val Gly Val Gly Ala Val 10 Ser Arg Asp Leu Glu Lys His Gly Ala Ile Thr Ser Ser Asn Thr Ala Ala Thr Asn Ala Asp Cys Ala Trp Leu Glu Ala Gln Glu Asp Glu Glu Val Gly Phe Pro Val Arg Pro Gln Val Pro Leu Arg Pro Met Thr Tyr Lys Gly Ala Val Asp Leu Ser His Phe Leu Lys Glu Lys Gly Gly Leu Glu Gly Leu Ile His Ser Gln Lys Arg Gln Asp Ile Leu Asp Leu Trp 15 Val Tyr His Thr Gln Gly Tyr Phe Pro Asp Trp Gln Asn Tyr Thr Pro Gly Pro Gly Ile Arg Phe Pro Leu Thr Phe Gly Trp Cys Phe Lys Leu Val Pro Val Glu Pro Glu Lys Val Glu Glu Ala Asn Glu Gly Glu Asn Asn Cys Leu Leu His Pro Met Ser Gln His Gly Ile Glu Asp Pro Glu Lys Glu Val Leu Glu Trp Arg Phe Asp Ser Lys Leu Ala Phe His His 20 Val Ala Arg Glu Leu His Pro Glu Tyr Tyr Lys Asp Cys (SEQ ID NO:12). Therefore, this exemplified Nef protein, Opt tPA-Nef, contains both a tPA leader sequence as well as deleting the myristylation site of Gly-2A DNA molecule encoding HIV-1 Nef from the HIV-1 jfrl isolate wherein the codons are optimized for expression in a mammalian system such as a human. 25

In another specific embodiment of the present invention, a DNA molecule is disclosed which encodes optimized HIV-1 Nef wherein the open reading frame of a recombinant adenoviral HIV vaccine encodes for modifications at the amino terminal myristylation site (Gly-2 to Ala-2) and substitution of the Leu-174-Leu-175 dileucine motif to Ala-174-Ala-175. This open reading frame is herein described as opt nef (G2A,LLAA) and is disclosed as SEQ ID NO:13, which comprises an initiating methionine residue at nucleotides 12-14 and a "TAA" stop codon from nucleotides 660-662. The nucleotide sequence of this codon optimized version of HIV-1 jrfl nef gene with the above mentioned modifications is disclosed herein as SEQ ID NO:13, as follows:

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The open reading frame of SEQ ID NO:13 encodes Nef (G2A,LLAA), disclosed herein as SEQ ID NO:14, as follows:

Met Ala Gly Lys Trp Ser Lys Arg Ser Val Pro Gly Trp Ser Thr Val 15 Arg Glu Arg Met Arg Arg Ala Glu Pro Ala Ala Asp Arg Val Arg Arg Thr Glu Pro Ala Ala Val Gly Val Gly Ala Val Ser Arg Asp Leu Glu Lys His Gly Ala Ile Thr Ser Ser Asn Thr Ala Ala Thr Asn Ala Asp Cys Ala Trp Leu Glu Ala Gln Glu Asp Glu Glu Val Gly Phe Pro Val Arg Pro Gln Val Pro Leu Arg Pro Met Thr Tyr Lys Gly Ala Val Asp 20 Leu Ser His Phe Leu Lys Glu Lys Gly Gly Leu Glu Gly Leu Ile His Ser Gln Lys Arg Gln Asp Ile Leu Asp Leu Trp Val Tyr His Thr Gln Gly Tyr Phe Pro Asp Trp Gln Asn Tyr Thr Pro Gly Pro Gly Ile Arg Phe Pro Leu Thr Phe Gly Trp Cys Phe Lys Leu Val Pro Val Glu Pro Glu Lys Val Glu Glu Ala Asn Glu Gly Glu Asn Asn Cys Ala Ala His 25 Pro Met Ser Gln His Gly Ile Glu Asp Pro Glu Lys Glu Val Leu Glu Trp Arg Phe Asp Ser Lys Leu Ala Phe His His Val Ala Arg Glu Leu His Pro Glu Tyr Tyr Lys Asp Cys Ser (SEQ ID NO:14).

An additional embodiment of the present invention relates to another DNA molecule encoding optimized HIV-1 Nef wherein the amino terminal myristylation site and dileucine motif have been deleted, as well as comprising a tPA leader peptide. This DNA molecule, opt tpanef (LLAA) comprises an open reading frame which encodes a Nef protein containing a tPA leader sequence fused to amino acid residue 6-216 of HIV-1 Nef (jfrl), wherein Leu-174 and Leu-175 are substituted with Ala-174 and Ala-175 (Ala-195 and Ala-196 in this tPA-based fusion protein). The nucleotide

sequence comprising the open reading frame of opt tpanef (LLAA) is disclosed herein as SEQ ID NO:15, as shown below:

CATGGATGCA ATGAAGAGA GGCTCTGCTG TGTGCTGCTG CTGTGTGGAG CAGTCTTCGT TTCGCCCAGC GAGATCTCCT CCAAGAGGTC CGTGCCCGGC TGGTCCACCG TGAGGGAGAG GATGAGGAGG GCCGAGCCCG CCGCCGACAG GGTGAGGAGG ACCGAGCCCG CCGCCGTGGG CGTGGGCGCC GTGTCCAGGG ACCTGGAGAA GCACGGCGCC ATCACCTCCT CCAACACCGC CGCCACCAAC GCCGACTGCG CCTGGCTGGA GGCCCAGGAG GACGAGGAGG TGGGCTTCCC CGTGAGGCCC CAGGTGCCCC TGAGGCCCAT GACCTACAAG GGCGCCGTGG ACCTGTCCCA CTTCCTGAAG GAGAAGGGCG GCCTGAGGGG CCTGATCCAC TCCCAGAAGA GGCAGGACAT 10 CCTGGACCTG TGGGTGTACC ACACCCAGGG CTACTTCCCC GACTGGCAGA ACTACACCCC CGGCCCGGC ATCAGGTTCC CCCTGACCTT CGGCTGGTGC TTCAAGCTGG TGCCCGTGGA GCCCGAGAAG GTGGAGGAGG CCAACGAGGG CGAGAACAAC TGCGCCGCCC ACCCCATGTC CCAGCACGC ATCGAGGACC CCGAGAAGGA GGTGCTGGAG TGGAGGTTCG ACTCCAAGCT GGCCTTCCAC CACGTGGCCA GGGAGCTGCA CCCCGAGTAC TACAAGGACT GCTAAAGCCC 15 (SEQ ID NO:15).

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The open reading frame of SEQ ID NO:7 encoding tPA-Nef (LLAA), disclosed herein as SEQ ID NO:16, is as follows:

Met Asp Ala Met Lys Arg Gly Leu Cys Cys Val Leu Leu Leu Cys Gly Ala Val Phe Val Ser Pro Ser Glu Ile Ser Ser Lys Arg Ser Val Pro 20 Gly Trp Ser Thr Val Arg Glu Arg Met Arg Arg Ala Glu Pro Ala Ala Asp Arg Val Arg Arg Thr Glu Pro Ala Ala Val Gly Val Gly Ala Val Ser Arg Asp Leu Glu Lys His Gly Ala Ile Thr Ser Ser Asn Thr Ala Ala Thr Asn Ala Asp Cys Ala Trp Leu Glu Ala Gln Glu Asp Glu Glu Val Gly Phe Pro Val Arg Pro Gln Val Pro Leu Arg Pro Met Thr Tyr 25 Lys Gly Ala Val Asp Leu Ser His Phe Leu Lys Glu Lys Gly Leu Glu Gly Leu Ile His Ser Gln Lys Arg Gln Asp Ile Leu Asp Leu Trp Val Tyr His Thr Gln Gly Tyr Phe Pro Asp Trp Gln Asn Tyr Thr Pro Gly Pro Gly Ile Arg Phe Pro Leu Thr Phe Gly Trp Cys Phe Lys Leu Val Pro Val Glu Pro Glu Lys Val Glu Glu Ala Asn Glu Gly Glu Asn Asn Cys Ala Ala His Pro Met Ser Gln His Gly Ile Glu Asp Pro Glu 30 Lys Glu Val Leu Glu Trp Arg Phe Asp Ser Lys Leu Ala Phe His His Val Ala Arg Glu Leu His Pro Glu Tyr Tyr Lys Asp Cys (SEQ ID NO:16). An adenoviral vector of the present invention may comprise a DNA sequence, regardless of codon usage, which expresses a wild type or modified Nef protein as described herein, including but not limited to modified Nef proteins which comprise a 35 deletion or substitution of Gly 2, a deletion of substitution of Leu 174 and Leu 175

and/or inclusion of a leader sequence. Therefore, partial or fully codon optimized DNA vaccine expression vector constructs are preferred since such constructs should result in increased host expression. However, it is within the scope of the present invention to utilize "non-codon optimized" versions of the constructs disclosed herein, especially modified versions of HIV Nef which are shown to promote a substantial cellular immune response subsequent to host administration.

Figure 20A-C show nucleotide sequences at junctions between nef coding sequence and plasmid backbone of nef expression vectors V1Jns/nef (Figure 20A), V1Jns/nef(G2A,LLAA) (Figure 20B), V1Jns/tpanef (Figure 20C) and V1Jns/tpanef(LLAA) (Figure 20C, also). 5' and 3' flanking sequences of codon optimized nef or codon optimized nef mutant genes are indicated by bold/italic letters; nef and nef mutant coding sequences are indicated by plain letters. Also indicated (as underlined) are the restriction endonuclease sites involved in construction of respective nef expression vectors. V1Jns/tpanef and V1Jns/tpanef(LLAA) have identical sequences at the junctions.

Figure 21 shows a schematic presentation of nef and nef derivatives. Amino acid residues involved in Nef derivatives are presented. Glycine 2 and Leucine 174 and 175 are the sites involved in myristylation and dileucine motif, respectively.

20 EXAMPLE 19

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MRKAd5Pol Construction and Virus Rescue

construction of vector: shuttle plasmid and pre-adenovirus plasmid - Key steps performed in the construction of the vectors, including the pre-adenovirus plasmid denoted MRKAd5pol, is depicted in Figure 22. Briefly, the adenoviral shuttle vector for the full-length inactivated HIV-1 pol gene is as follows. The vector MRKpdelE1(Pac/pIX/pack450)+CMVmin+BGHpA(str.) is a derivative of the shuttle vector used in the construction of the MRKAd5gag adenoviral pre-plasmid. The vector contains an expression cassette with the hCMV promoter (no intronA) and the bovine growth hormone polyadenylation signal. The expression unit has been inserted into the shuttle vector such that insertion of the gene of choice at a unique BgIII site will ensure the direction of transcription of the transgene will be Ad5 E1 parallel when inserted into the MRKpAd5(E1-/E3+)Cla1 (or MRKpAdHVE3) pre-plasmid. The vector, similar to the original shuttle vector contains the Pac1 site, extension to the packaging signal region, and extension to the pIX gene. The synthetic full-length codon-optimized HIV-1 pol gene was isolated directly from the plasmid pV1Jns-HIV-pol-inact(opt). Digestion of this plasmid with BgI II releases the pol

gene intact (comprising a codon optimized IA pol sequence as disclosed in SEQ ID NO:3). The pol fragment was gel purified and ligated into the MRKpdelE1(Pac/pIX/pack450)+CMVmin+BGHpA(str.) shuttle vector at the BgIII site. The clones were checked for the correct orientation of the gene by using restriction enzymes Dra III/Not1. A positive clone was isolated and named MRKpdel+hCMVmin+FL-pol+bGHpA(s). The genetic structure of this plasmid was verified by PCR, restriction enzyme and DNA sequencing. The pre-adenovirus plasmid was constructed as follows. Shuttle plasmid MRKpdel+hCMVmin+FLpol+bGHpA(S) was digested with restriction enzymes Pac1 and Bst1107 I (or its isoschizomer, BstZ107 I) and then co-transformed into E. coli strain BJ5183 with linearized (Cla1 digested) adenoviral backbone plasmid, MRKpAd(E1-/E3+)Cla1. The resulting pre-plasmid originally named MRKpAd+hCMVmin+FLpol+bGHpA(S)E3+ is now referred to as "pMRKAd5pol". The genetic structure of the resulting pMRKAd5pol was verified by PCR, restriction enzyme and DNA sequence analysis. The vectors were transformed into competent E. coli XL-1 Blue for preparative production. The recovered plasmid was verified by restriction enzyme digestion and DNA sequence analysis, and by expression of the pol transgene in transient transfection cell culture. The complete nucleotide sequence of this pMRKAd5HIV-1pol adenoviral vector is shown in Figure 26 A-AO.

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Generation of research-grade recombinant adenovirus - The pre-adenovirus plasmid, pMRKAd5pol, was rescued as infectious virions in PER.C6® adherent monolayer cell culture. To rescue infectious virus, 12 μ g of pMRKAd5pol was digested with restriction enzyme Pacl (New England Biolabs) and 3.3 μ g was transfected per 6 cm dish of PER.C6® cells using the calcium phosphate coprecipitation technique (Cell Phect Transfection Kit, Amersham Pharmacia Biotech Inc.). Pacl digestion releases the viral genome from plasmid sequences allowing viral replication to occur after entry into PER.C6® cells. Infected cells and media were harvested 6 -10 days post-transfection, after complete viral cytopathic effect (CPE) was observed. Infected cells and media were stored at \leq -60°C. This pol containing recombinant adenovirus is referred to herein as "MRKAd5pol". This recombinant adenovirus expresses an inactivated HIV-1 Pol protein as shown in SEQ ID NO:6.

EXAMPLE 20

MRKAd5Nef Construction and Virus Rescue

Construction of vector: shuttle plasmid and pre-adenovirus plasmid - Key steps performed in the construction of the vectors, including the pre-adenovirus

plasmid denoted MRKAd5nef, is depicted in Figure 23. Briefly, as shown in Example 19 above, the vector

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MRKpdelE1(Pac/pIX/pack450)+CMVmin+BGHpA(str.) is the shuttle vector used in the construction of the MRKAd5gag adenoviral pre-plasmid. It has been modified to contain the *Pac*1 site, extension to the packaging signal region, and extension to the pIX gene. It contains an expression cassette with the hCMV promoter (no intronA) and the bovine growth hormone polyadenylation signal. The expression unit has been inserted into the shuttle vector such that insertion of the gene of choice at a unique *Bgl*11 site will ensure the direction of transcription of the transgene will be Ad5 E1 parallel when inserted into the MRKpAd5(E1-/E3+)Cla1 pre-plasmid. The synthetic full-length codon-optimized HIV-1 nef gene was isolated directly from the plasmid pV1Jns/nef (G2A,LLAA). Digestion of this plasmid with *Bgl*11 releases the pol gene intact, which comprises the nucleotide sequence as disclosed in SEQ ID NO:13. The nef fragment was gel purified and ligated into the

MRKpdelE1+CMVmin+BGHpA(str.) shuttle vector at the *Bgl*11 site. The clones were checked for correction orientation of the gene by using restriction enzyme *Sca*1. A positive clone was isolated and named MRKpdelE1hCMVminFL-nefBGHpA(s). The genetic structure of this plasmid was verified by PCR, restriction enzyme and DNA sequencing. The pre-adenovirus plasmid was constructed as follows. Shuttle plasmid MRKpdelE1hCMVminFL-nefBGHpA(s) was digested with restriction enzymes *Pac*1 and *Bst*1107 I (or its isoschizomer, *Bst*Z107 I) and then co-transformed into *E. coli* strain BJ5183 with linearized (*Cla*1 digested) adenoviral backbone plasmid, MRKpAd(E1/E3+)Cla1. The resulting pre-plasmid originally named MRKpdelE1hCMVminFL-nefBGHpA(s) is now referred to as "pMRKAd5nef". The genetic structure of the resulting pMRKAd5nef was verified by PCR, restriction enzyme and DNA sequence analysis. The vectors were transformed into competent *E. coli* XL-1 Blue for preparative production. The recovered plasmid was verified by restriction enzyme digestion and DNA sequence analysis, and by expression of the nef transgene in transient transfection cell culture. The complete nucleotide sequence

Generation of research-grade recombinant adenovirus - The pre-adenovirus plasmid, pMRKAd5nef, was rescued as infectious virions in PER.C6® adherent monolayer cell culture. To rescue infectious virus, 12 μ g of pMRKAdnef was digested with restriction enzyme Pac1 (New England Biolabs) and 3.3 μ g was transfected per 6 cm dish of PER.C6® cells using the calcium phosphate coprecipitation technique (Cell Phect Transfection Kit, Amersham Pharmacia Biotech

of this pMRKAd5HIV-1nef adenoviral vector is shown in Figure 27A-AM.

Inc.). Pac1 digestion releases the viral genome from plasmid sequences allowing viral replication to occur after entry into PER.C6[®] cells. Infected cells and media were harvested 6-10 days post-transfection, after complete viral cytopathic effect (CPE) was observed. Infected cells and media were stored at \leq -60°C. This nef containing recombinant adenovirus is now referred to as "MRKAd5nef".

EXAMPLE 21

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Construction of Murine CMV Promoter Containing Shuttle Vectors for Inactivated Pol and Nef/G2A,LLAA

The murine CMV (mCMV) was amplified from the plasmid pMH4 (supplied 10 by Frank Graham, McMaster University) using the primer set: mCMV (Not I) Forward: 5'-ATA AGA ATG CGG CCG CCA TAT ACT GAG TCA TTA GG-3' (SEO ID NO: 20); mCMV (Bgl II)Reverse: 5'-AAG GAA GAT CTA CCG ACG CTG GTC GCG CCT C-3' (SEQ ID NO:21). The underlined nucleotides represent the Not I and the $Bgl \Pi$ sites respectively for each primer. This PCR amplicon was 15 used for the construction of the mCMV shuttle vector containing the transgene in the El parallel orientation. The hCMV promoter was removed from the original shuttle vector (containing the hCMV-gag-bGHpA transgene in the E1 parallel orientation) by digestion with Not I and Bgl II. The mCMV promoter (Not I/Bgl II digested PCR product) was inserted into the shuttle vector in a directional manner. The shuttle 20 vector was then digested with $Bgl \Pi$ and the gag reporter gene ($Bgl \Pi$ fragment) was re-inserted back into the shuttle vector. Several clones were screened for correct orientation of the reporter gene. For the construction of the mCMV-gag in the E1 antiparallel orientation, the mCMV promoter was amplified from the plasmid pMH4 using the following primer set: mCMV (Asc I) Forward: 5'- ATA AGA ATG GCG 25 CGC CAT ATA CTG AGT CAT TAG G (SEQ ID NO:22); mCMV (Bgl II) Reverse: 5' AAG GAA GAT CTA CCG ACG CTG GTC GCG CCT C (SEQ ID NO:23). The underlined nucleotides represent the Asc I and Bgl II sites, respectively for each primer. The shuttle vector containing the hCMV-gag transgene in the E1 antiparallel orientation was digested with Asc1 and Bgl11 to remove the hCMV-gag portion of the 30 transgene. The mCMV promoter (Asc1/Bgl11 digested PCR product) was inserted into the shuttle vector in a directional manner. The vector was then digested with Bgl11 and the gag reporter gene (Bgl11 fragment) was re-inserted. Several clones were screened for correct orientation of the reporter gene. For each of the full length IA pol and full length nef/G2A,LLAA genes, cloning was performed using the unique 35

 $Bgl~\Pi$ site within the mCMV-bGHpA shuttle vector. The pol and nef genes were excised from their respective pV1Ins plasmids by $Bgl~\Pi$ digestion.

EXAMPLE 22

Construction of mCMV Full Length Inactivated Pol and Full Length nef/G2A.LLAA Adenovectors

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Each of these transgenes of Example 21 were inserted into the modified shuttle vector in both the E1 parallel and E1 anti-parallel orientations. *Pac1* and *BstZ110I* digestion of each shuttle vector was performed and each specific transgene fragment containing the flanking Ad5 sequences was isolated and co-transformed with *Cla I* digested MRKpAd5(E3+) or MRKpAd5(E3-) adenovector plasmids via bacterial homologous recombination in BJ5183 *E. coli* cells. Recombinant preplasmid adenovectors containing the various transgenes in both the E3- and E3+ versions (and in the E1 parallel and E1 antiparallel orientations) were subsequently prepared in large scale following transformation into XL-1 Blue *E. coli* cells and analyzed by restriction analysis and sequencing.

EXAMPLE 23

Construction of hCMV-tpa-nef (LLAA) Adenovector

The tpa-nef gene was amplified out from GMP grade pV1Jns-tpanef (LLAA) vector using the primer sets: Tpanef (BamHI) F 5'-ATT GGA TCC ATG GAT GCA ATG AAG AGA GGG (SEQ ID 24); Tpanef (BamHI) R 5'-ATA GGA TCC TTA GCA GTC CTT GTA GTA CTC G (SEQ ID NO:25). The resulting PCR product was digested with BamHI, gel purified and cloned into the Bgl II site of MRKAd5CMV-bGHpA shuttle vector (Bgl II digested and calf intestinal phosphatase treated). Clones containing the tpanef (LLAA) gene (see SEQ ID NO:15 for complet coding region) in the correct orientation with respect to the hCMV promoter were selected following Sca I digestion. The resulting MRKAd5tpanef shuttle vector was digested with Pac I and Bst Z1101 and cloned into the E3+ MRKAd5 adenovector via bacterial homologous recombination techniques.

EXAMPLE 24

Immunogenicity of MRKAd5pol and MRKAd5nef Vaccine

Materials and Methods - Rodent Immunization - Groups of N=10 BALB/c

mice were immunized i.m. with the following vectors: (1) MRKAd5hCMV-IApol

(E3+) at either 10^7 vp and 10^9 vp; and (2) MRKAd5hCMV-IApol (E3-) at either

10^7 vp and 10^9 vp. At 7 weeks post dose, 5 of the 10 mice per cohort were boosted with the same vector and dose they initially received. At 3 weeks post the second does, sera and spleens were collected from all the animals for RT ELISA and IFNg ELIspot analyses, respectively. For all rodent immunizations, the Ad5 vectors were diluted in 5 mM Tris, 5% sucrose, 75 mM NaCl, 1 mM MgCl2, 0.005% polysorbate 80, pH 8.0. The total dose was injected to both quadricep muscles in 50 µL aliquots using a 0.3-mL insulin syringe with 28-1/2G needles (Becton-Dickinson, Franklin Lakes, NJ).

Groups of N=10 C57/BL6 mice were immunized i.m. with the following vectors: (1) MRKAd5hCMV-nef(G2A,LLAA) (E3+) at either 10^7 vp and 10^9 vp; (2) MRKAd5mCMV-nef(G2A,LLAA) (E3+) at either 10^7 vp and 10^9 vp; and (3) MRKAd5mCMV-tpanef(LLAA) (E3+) at either 10^7 vp and 10^9 vp. At 7 weeks post dose, 5 of the 10 mice per cohort were boosted with the same vector and dose they initially received. At 3 weeks post the second does, sera and spleens were collected from all the animals for RT ELISA and IFNg ELIspot analyses, respectively.

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Non-human Primate immunization - Cohorts of 3 rhesus macaques (2-3 kg) were vaccinated with the following Ad vectors: (1) MRKAd5hCMV-IApol (E3+) at either 10^9 vp and 10^11 vp dose; and (2) MRKAd5hCMV-IApol (E3-) at either 10^9 vp and 10^11 vp; (3) MRKAd5hCMV-nef(G2A,LLAA) (E3+) at either 10^9 vp and 10^11 vp; and (4) MRKAd5mCMV-nef(G2A,LLAA) (E3+) at either 10^9 vp and 10^11 vp. The vaccine was administered to chemically restrained monkeys (10 mg/kg ketamine) by needle injection of two 0.5 mL aliquots of the Ad vectors (in 5 mM Tris, 5% sucrose, 75 mM NaCl, 1 mM MgCl₂, 0.005% polysorbate 80, pH 8.0) into both deltoid muscles. The animals were immunized twice at a 4 week interval (T=0, 4 weeks).

Murine anti-RT and anti-nef ELISA - Anti-RT titers were obtained following standard secondary antibody-based ELISA. Maxisorp plates (NUNC, Rochester; NY) were coated by overnight incubation with 100 μL of 1 μg/mL HIV-1 RT protein (Advanced Biotechnologies, Columbia, MD) in PBS. For anti-nef ELISA, 100 uL of 1 ug/mL HIV-1 nef (Advanced Biotechnologies, Columbia, MD) was used to coat the plates. The plates were washed with PBS/0.05% Tween 20 using Titertek MAP instrument (Hunstville, AL) and incubated for 2 h with 200 μL/well of blocking solution (PBS/0.05% tween/1% BSA). An initial serum dilution of 100-fold was performed followed by 4-fold serial dilution. 100-μL aliquots of serially diluted samples were added per well and incubated for 2 h at room temperature. The plates

were washed and 100 μL of 1/1000-diluted HRP-rabbit anti-mouse IgG (ZYMED, San Francisco, CA) were added with 1 h incubation. The plates were washed thoroughly and soaked with 100 μL 1,2-phenylenediamine dihydrochloride/hydrogen peroxide (DAKO, Norway) solution for 15 min. The reaction was quenched by adding 100 μL of 0.5M H₂SO4 per well. OD₄₉₂ readings were recorded using Titertek Multiskan MCC/340 with S20 stacker. Endpoint titers were defined as the highest serum dilution that resulted in an absorbance value of greater than or equal to 0.1 OD₄₉₂ (2.5 times the background value).

Non-human primate and murine ELIspot assays - The enzyme-linked immuno-spot (ELISpot) assay was utilized to enumerate antigen-specific INFy-10 secreting cells from mouse spleens (Miyahira, et al. 1995, J. Immunol. Methods 181:45-54) or macaque PBMCs. Mouse spleens were pooled from 5 mice/cohort and single cell suspensions were prepared at 5x106/mL in complete RPMI media (RPMI1640, 10% FBS, 2mM L-glutamine, 100U/mL Penicillin, 100 u/mL streptomycin, 10 mM Hepes, 50 uM β-ME). Rhesus PBMCs were prepared from 8-15 15 mL of heparinized blood following standard Ficoll gradient separation (Coligan, et al, 1998, Current Protocols in Immunology. John Wiley & Sons, Inc.). Multiscreen opaque plates (Millipore, France) were coated with 100 μL/well of either 5 μg/mL purified rat anti-mouse IFN-y IgG1, clone R4-6A2 (Pharmingen, San Diego, CA), or 15 ug/mL mouse anti-human IFN-γ IgG_{2a} (Cat. No. 1598-00, R&D Systems, 20 Minneapolis, MN) in PBS at 4°C overnight for murine or monkey assays, respectively. The plates were washed with PBS/penicillin/streptomycin and blocked with 200 μL/well of complete RPMI media for 37 °C for at least 2 h.

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To each well, 50 μL of cell samples (4-5x10⁵ cells per well) and 50 μL of the antigen solution were added. To the control well, 50 μL of the media containing DMSO were added; for specific responses, either selected peptides or peptide pools (4 ug/mL per peptide final concentration) were added. For BALB/c mice immunized with the pol constructs, stimulation was conducted using a pool of CD4⁺-epitope containing 20-mer peptides (aa21-40, aa411-430, aa641-660, aa731-750, aa771-790) or a pool of CD8⁺-epitope containing peptides (aa201-220, aa311-330, aa781-800). For C57/BL6 mice immunized with the nef construct, either aa51-70 (CD8⁺ T cell epitope) or aa81-100 (CD4⁺) peptide derived from the nef sequence was added for specific stimulation. In monkeys, the responses against pol were evaluated using two pools (L and R) of 20-aa peptides that encompass the entire pol sequence and overlap by 10 amino acids. In monkeys vaccinated with the nef constructs, a single pool containing 20-mer peptides covering the entire HIV-1 nef sequence and overlapping

by 10 aa was used. Each sample/antigen mixture was performed in triplicate wells for murine samples or in duplicate wells for rhesus PBMCs. Plates were incubated at 37°C, 5% CO₂, 90% humidity for 20-24 h. The plates were washed with PBS/0.05% Tween 20 and incubated with 100 μL/well of either 1.25 μg/mL biotin-conjugated rat anti-mouse IFN-γ mAb, clone XMG1.2 (Pharmingen) or of 0.1 ug/mL biotinylated anti-human IFN-gamma goat polyclonal antibody (R&D Systems) at 4°C overnight. The plates were washed and incubated with 100 μL/well 1/2500 dilution of strepavidin-alkaline phosphatase conjugate (Pharmingen) in PBS/0.005% Tween/5% FBS for 30 min at 37 °C. Spots were developed by incubating with 100 μL/well 1-step NBT/BCIP (Pierce Chemicals) for 6-10 min. The plates were washed with water and allowed to air dry. The number of spots in each well was determined using a dissecting microscope and the data normalized to 10⁶ cell input.

Non-human Primate anti-RT ELISA - The pol-specific antibodies in the monkeys were measured in a competitive RT EIA assay, wherein sample activity is determined by the ability to block RT antigen from binding to coating antibody on the plate well. Briefly, Maxisorp plates were coated with saturating amounts of pol positive human serum (#97111234). 250 uL of each sample is incubated with 15 uL of 266 ng/mL RT recombinant protein (in RCM 563, 1% BSA, 0.1% tween, 0.1% NaN₃) and 20 uL of lysis buffer (Coulter p24 antigen assay kit) for 15 min at room temperature. Similar mixtures are prepared using serially diluted samples of a standard and a negative control which defines maximum RT binding. 200 uL/well of each sample and standard were added to the washed plate and the plate incubated 16-24 h at room temperature. Bound RT is quantified following the procedures described in Coulter p24 assay kit and reported in milliMerck units per mL arbitrarily defined by the chosen standard.

Results - Rodent Studies - BALB/c mice (n=5 mice/cohort) were immunized once or twice with varying doses of MRKAd5hCMV-IApol(E3+) and MRKAd5hCMV-IApol(E3-). At 3 weeks after the second dose, Anti-pol IgG levels were determined by an ELISA assay using RT as a surrogate antigen. Cellular response were quantified via IFNY ELISpot assay against pools of pol-epitope containing peptides. The results of these assays are summarized in Table 10. The results indicate that the mouse vaccinees exhibited detectable anti-RT IgGs with an adenovector dose as low as 10^7 vp. The humoral responses are highly dose-dependent and are boostable with a second immunization. One or two doses of either pol vectors elicit high frequencies of antigen-specific CD4⁺ and CD8⁺ T cells; the responses are weakly dose-dependent but are boostable with a second immunization.

Table 10. Immunogenicity of MRKAd5pol Vectors in BALB/c mice.

	20.			An	II-RT InG Tite	rs"	S	FC/10^6 cell	s°_
Group	Vaccine	Dose	No. of Doses	GMT	+SE	-SE	Medium	CD4+ peptide pool	CD8+ peptide pool
1	MRKAd5hCMVFLpol (E3+)	10^7 vp	2 1	310419 919	301785 372	153020 265	1(1) 1(1)	75(4) 72(9)	2313(67) 533(41)
2	MRKAd5hCMVFLpol (E3+)	10^9 vp	2	1638400 ⁶ 713155	0 528520	0 303555	2(2) 1(1)	114(9) 48(7)	2063(182) 733(89)
3	MRKAd5hCMVFLpol (E3-)	10^7 vp	2	310419 6400	386218 14013	172097 4393	0(0) 10(8)	223(7) 141(21)	2607(27) 409(28)
4	MRKAd5hCMVFLpol (E3-)	10^9 vp	2	1638400 ^b 1241675 ^b	0 396725	0 300681	1(1) 0(0)	160(13) 39(13)	2385(11) 833(83)
5	Naīve	none	none	57	9	7	9(2)	11(4)	10(1)

^{*}GMT, geometric mean titer of the cohort of 5 mice; SE, standard error of the gemetric mean

C57/BL6 mice were immunized once or twice with varying doses of MRKAd5hCMV-nef(G2A,LLAA) (E3+), MRKAd5mCMV-nef(G2A,LLAA) (E3+) at either 10^7 vp and(3) MRKAd5mCMV-tpanef(LLAA) (E3+) at either 10^7 vp and 10^9 vp. The immune response were analyzed using similar protocols and the results are listed in Table 11. While anti-nef IgG responses could not be detected in this model system with any of the constructs, there are strong indications of a cellular immunity generated against nef using the ELIspot assay.

Table 11 Immunogenicity of MRKAd5nef Vectors in C57/BL6 mice.

				An	ti-nef IgG Tite	rs	s	FC/10^6 cell	
Group	Vaccine	Dose	No. of Doses	GMT	+SE	-SE	Medium	aa51-70 CD8+	aa81-100 CD4+
1	MRKAd5hCMVFLnef (E3+)	10^7 vp	2 1	174 132	70 42	50 32	1(1) 0(0)	23(1) 0(0)	1(1) 0(0)
2	MRKAd5hCMVFLnef (E3+)	10 ^9 vp	2	174 132	70 42	50 32	0(0) 1(1)	61(7) 62(7)	4(2) 3(1)
3	MRKAd5mCMVFLnef (E3+)	10^7 vp	2 1 ·	132 115	42 46	32 33	3(1) 3(2)	15(5) 3(2)	5(2) 4(2)
4	MRKAd5mCMVFLnel (E3+)	10^9 vp	2	132 132	42 42	32 32	4(2) 2(1)	83(13) 29(2)	5(1) 4(0)
5	MRKAd5mCMVtpanef(E3+)	10^7 vp	2	132 100	42 0	32	3(2) 3(1)	14(2) 13(4)	5(1) 10(3)
6	MRKAd5mCMVtpanef(E3+)	10^9 vp	2	230 115	170 46	98 33	3(2) 7(1)	145(29) 151(14)	4(0) 10(0)
	Naīve	none	none	152	78	52	· 21(2)	- 18(6)	28(3)

^{*}GMT, geometric mean ther of the cohort of 5 mice; SE, standard error of the gemetric mean

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Monkey Studies - Cohorts of 3 rhesus macaques were immunized with 2 doses of MRKAd5hCMV-IApol(E3+) and MRKAd5hCMV-IApol(E3-). The number of antigen-specific T cells (per million PBMCs) were enumerated using one of two

Near or at the upper limit of the serial dilution; hence, could be greater than this value

^{*}No. of Spot-forming Cells per million splechoytes; mean values of triplicates are reported along with standard errors in parenthesis.

No. of spot-forming cells per million splecnoyles; mean values of triplicates are reported along with standard errors in parenthesis.

peptide pools (L and R) that cover the entire pol sequence; the results are listed in Table 12. Moderate-to-strong T cell responses were detected in the vaccinees using either constructs even at a low dose of 10^9 vp. Longitudinal analyses of the anti-RT antibody titers in the animals suggest that the pol transgene product is expressed efficiently to elicit a humoral response (Table 13). It would appear that generally higher immune responses were observed in animals that received the E3- construct compared to the E3+ virus.

Table 12. Pol-specific T Cell Responses in MRKAd5pol Immunized Rhesus

10 Macaques.

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Vaccine (T=0,4 wks)	Monk #		Prebleed			T=4			T=7			T=16	
		Mock	Pol L	Pol R	Mock	Pol L	Pol R	Mock	PolL	Pol R	Mock	Pal L	Pol R
MRKAd5hCMV-IApol(E3+)	99C100	1	0	0	1	38	31	0	52	146	0	49	715
10413 AD	99C215	1	2	2	10	98	249	1 1	109	305	22	88	250
10°11 VP	99D201	5	5	4	6	149	85	0	40	35	0	35	18
MRKACEHOMV-IApol(E3+)	99D212	0	2	0	4	331	114	0	58	14	0	6	6
10/9 VP	99D180	0	4	2	0	19	192	4	38	158	5	38	108
	99C201	8	5	21	6	82	62	0	18	32	١ ١	14	65
MRKAdShCMV-IApol(E3-)	99D239	5	2	2	20	82	172	1	68	114	9	21	40
10^11 VD	99C186	4	12	6	5	120	421	2	271	489	16	875	530
	99C084	1	8	8	8	84	484	0	14	238	١	24	264
MRKAd5hCMV-IApol(E3-)	∞ 7с	10	10	8	12	724	745	4	322	376	4	188	176
10/9 VD	CDIG	2	0	1 1	5	474	468	0	232	212	0	101	121
10 / 1 p	CD11	6	6	12	10	98	110	5	60	80	8	25	34
Nave	083Q	nd	nd	nd	nd	nd	nd	4	2	2	2	1_1_	2

nd, not determined Reported are SFC per million PBMCs; mean of duplicate wells.

Table 13. Anti-RT Ig Levels in MRKAd5pol Immunized macaques.

RT ANTIBODY ASSAYTITERS IN MMU/	mL			
Vaccine/Monkey Tag	1=4	T=7	T=12	T=16
MRKAd5hCMV-IApol(E3+), 10^11 vp				
99C100	61	1999	5928	4768
99C215	81	1541	2356	2767
99D201	53	336	539	387
MRKAd5hCMV-IApol(E3+), 10^9 vp			·	
99D212	10	40	49	68
99D180	<10	36	79	93
99C201	<10	37	71	76
MRKAd5hCMV-IApol(E3-), 10^11 vp				
99D239	44	460	1234	1015
99C186	21	· 233 ·	480	345
990084	235	2637	2858_	1626
MRKAd5hCMV-IApol(E3-), 10^9 vp				
CC7C	32	175	306	235
ωie	20	140	273	419
Q11	15	112	149	237

When rhesus macaques were immunized i.m. with two doses of MRKAd5nef constructs, vigorous T cell responses ranging from 100 to as high as 1100 per million were observed in 8 of 12 vaccinees (Table 14). The efficacies of the mCMV- and hCMV- driven nef constructs are comparable on the basis of the data generated thus far.

Table 14. Nef-specific T cell Responses in MRKAd5nef Immunized Rhesus

Macaques.

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Vaccine (T=0,4 wks)	Monk#	Pr	e	Te	4	T=	.7	T≃	16
		Mock	Nef	Mock	Nef	Mock	Net	Mock	Nef
MRKAd5hCMV-nef(G2A,LLAA) (E3+)	CD2D	0	4	31	440	4	368	1	251
10^11 vp	CC7B	1 0 1	0	2	521	0	178	1	152
	CC61	2	9	31	112	0	108	11	100
MRKAd5hCMV-net(G2A,LLAA) (E3+)	CC2K	9	9	6	52	0	35	0	15
10^9 vp	CD15	5	4	30	998	2	586	0	434
	CD16	6	1	6	1146	0	369	1	212
MRKAd5mCMV-nef(G2A,LLAA) (E3+)	99D191	1	5	4	614	0	298	2	418
10^11 vp	99D144	4	6	5	434	0	1100	2	933
	99C193	1 1	2	1	58	1	22	0	64
MRKAd5mCMV-nef(G2A,LLAA) (E3+)	99D224	1	11	14	231	1	125	0	70
10^9 VD	99D250	8	9	4	108	0	54	0	5
	99C120	1	6	20	299	0	92	٥	75
Naîve	083Q	nd	nd	18	22	4	5	2	1

EXAMPLE 25

Comparison of Clade B vs. Clade C T Cell Responses in HIV-Infected Subjects
PBMC samples collected from two dozens of patients infected with HIV-1 in
US were tested in ELISPOT assays with peptide pools of 20-mer peptides overlapping
by 10 amino acids. Four different peptide pools were tested for cross-clade
recognition, and they were either derived from a clade B-based isolate (gag H-b; nefb) or a clade C-based isolate (gag H-c, nef-c). Data in Table 15 shows that T cells
from these patients presumably infected with clade B HIV-1 could recognize clade C
gag and nef antigens in ELISPOT assay. Correlation analysis further demonstrated
that these T cell responses against clade C gag peptide pool were about 60% of the
clade B counterpart (Figure 24), while the T cell responses against clade C nef were
about 85% of the clade B counterpart (Figure 25). These results suggest that cellular
immune responses generated in patients infected with clade B HIV-1 can recognize
gag and nef antigens derived from clade C HIV-1. These data show that a HIV
vaccine, such as a DNA or MRKAd5-based adenoviral vaccine expressing a clade B

gag and/or nef antigen will potentially have the ability to provide a prophylactic and/or therapetic advantage on a global scale.

Table 15
Responses Shown as the Number of gIFN-Secreting T Cells per Million PBMCs

subject	bleed date	gag epitope #	mock	gag H-b	gagH-c	nef-b	nef-c
		from mapping)					
#100	19-Jul-99	12	10	3950	1385	1295	1300
#101	25-Jul-99	3	15	3885	1280	na	1020
#102	25-Jul-99	4	15	1740	850	1255	1785
#104	7-Jun-99	2	5	1355	1185	na	1060
#107	11-Oct-99	2	25	3305	2795	670	870
#405	11-Jul-99	2	15	4575	3180	1700	1500
#501	19-Jul-99	2	15	1100	570	3365	3460
#505	18-Jul-99	5	10	2145	1725	1235	na
#506	28-Feb-99	2	25	150	45	400	610
#701	28-Mar-99	5	30	7620	4775	3320	2780
#709	17-May-99	3	15	2785	1945	1090	1630
#710	24-May-99	4	5	1055	1080	2210	2140
				1			

10 EXAMPLE 26

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Characterization and Production of MRKAd5pol and MRKAd5nef Vectors in Roller Bottles

Expansion of nef and pol Adenovectors - Nef and pol CsCl purified MRKAd5 seeds were used to infect roller bottles to produce P4 virus to be used as a seed for further experiments. P4 MRKAd5 pol and nef vectors were used to infect roller bottles at an MOI 280 vp/cell, except for hCMV-tpa-nef [E3+] which was infected at an MOI of 125 due to low titers of seed obtained at P4.

Table 16 Viral particle concentrations for P5 nef and pol adenovectors

Adenovector	AEX Titer (10 ¹⁰ vp/ml culture)	AEX Titer (10 ⁴ vp/cell)	Amplification Ratio
hCMV-FL-nef [E3+]	1.1	0.9	30
mCMV-FL-nef [E3+]	2.2	2.1	75
hCMV-tpa-nef [E3+]	0.07	0.1	5
mCMV-tpa-nef [E3+]	1.3	0.9	35
hCMV-FL-pol [E3+]	2.7	2.1	75
hCMV-FL-pol [E3-]	1.9	1.3	45

Roller Bottle Passaging - Passaging of the pol and nef constructs continued through passage seven. Cell-associated (freeze/thaw lysis) and whole broth (tritonlysis) titers obtained in all passages were very consistent. In general, MRKAd5pol is ca. 70% as productive as MRKAd5gag while MRKAd5nef is ca. 25% as productive as MRKAd5gag. Samples of P7 virus for both constructs were analyzed by V&CB by restriction digest analysis and did not show any rearrangements.

Table 17. Passage Six Viral Productivity for MRKAd5pol and MRKAd5nef

24010]	Xviable (10	oells/ml),	Cell Passage	AEX Titer	Titer	Amplification	Triton Lysis Titer
		Viabil Infection	ity (%) Harvest	Number	(Cell Associated) 10 ¹⁰ vp/ml culture	10 ⁴ vp/ccil	Ratio	10 ¹⁰ vp/ml culture
hCMV-FL-nef [B3+]	pool	1.22, 85%		62	0.8	0.7	25	1.6
	-1		0.99, 62%	1				
	2		1.10, 72%	1	}			
bCMV-FL-pol (E3+)	pool	1.42, 89%		62	4.5	3.2	115	7.0
	. 1		1.22, 70%					
	2		1.42, 74%					

15 Table 18. Passage Seven Viral Productivity for MRKAd5pol and MRKAd5nef

		Xviable (10 Viabil Infection	of cells/ml), ity (%) Harvest	Cell Passage Number	AEX Titer (Cell Associated) 10 ⁸⁰ vp/ml culture	Titer 10 ⁴ vp/cell	Amplification Ratio	Triton Lysis Titer 10 ¹⁰ vp/ml culture
hCMV-FL-ncf [E3+]	Pool	1.33, 90%		66	1.0	0.8	29	2.1
	1		0.96, 70%					
	2		1.18, 73%	·Ì				
hCMV-FL-pol [E3+]	Pool	0.90*, 90%		56	4.2	4.7	168	6.5
			1.18, 88%	1				
	2		1.04, 80%					

MRKAd5nef and MRKAd5pol Viral Production Kinetics - A timecourse experiment was carried out in roller bottles to determine if the viral production kinetics of the MRKAd5pol and MRKAd5nef vectors were similar to those of MRKAd5gag. PER.C6® cells in roller bottle cultures were infected at an MOI of 280 vp/cells with P5 MRKAd5pol, P5 MRKAd5nef and P7 MRKAd5gag; for each adenovector, two infected bottles were sampled at 24, 36, 48, and 60 hours post infection. In addition, two bottles were left unsampled until 48 hpi when they were harvested under the Phase I process conditions. The anion-exchange HPLC viral particle concentrations of the freeze-thaw recovered cell associated virus at the 24, 36,

48, and 60 hpi timepoints are shown in Figure 29A-B. The QPA titers show a similar trend (data not shown).

Comparison of hCMV- and mCMV-FL-nef - As the titers obtained with the MRKAd5nef construct (hCMV-FL-nef) were lower than those obtained with MRKAd5gag or MRKAd5pol, a viral productivity comparison experiment was performed with mCMV-FL-nef. For each of the two adenovectors (hCMV- and mCMV-FL-nef), two roller bottles were infected at an MOI of 280 vp/cell with passage five clarified lysate. The macroscopic and microscopic observations of the four roller bottles were identical at the time of harvest. Analysis of the clarified lysate produced indicated a higher viral particle concentration in the bottles infected with mCMV-FL-nef, as shown in Table 19. It is stipulated that the higher productivity with mCMV promoter driven nef vector is due to lower nef expression levels in PER.C6[®] cells- experiments are underway at V&CB to measure nef expression levels.

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Table 19. Passage Six Viral Productivity Comparison of hCMV- and mCMV-FL-nef

	[Xv (10° cells/m	l), Viability (%)	Cell Passage	AEX Titer	Titer	Amplification	Triton Lysis Titer
		Infection	Harvest	Number	10 ¹⁰ vp/ml culture	104 vp/cell	Ratio	10 ^{to} vp/ml culture
hCMV-FL-nef	Pool	1.11, 91%		60	1.5	1.4	50	2.8
(MRKAd5nef)	. 1		1.23, 75%				1	
	2		1.34,74%		1			
mCMV-FL-nef	Pool	1.11, 91%		60	2.3	2.1	75	4.6
	1		1.49, 84%		}			
	2		1.18, 77%					

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EXAMPLE 27

Characterization and Large Scale Production of MRKAd5nef Virus in Bioreactors

under the following conditions: 36.5°C, DO 30%, pH 7.30, 150rpm agitation rate, no sparging, Life Technologies (Gibco, Invitrogen) 293 SFM II (with 6mM L-glutamine), 0.5M NaOH as base for pH control. During the first run (B20010115), two 10L stirred vessel bioreactors were inoculated with PER.C6® cells at a concentration of 0.2x10° cells/ml. Cells were grown until they reached a cell concentration of approximately 1x10° cells/ml. The cells were infected with uncloned MRKAd5nef (G2A,LLAA) at a MOI of 280 virus particles (vp)/cell. For the second batch (B20010202), the same procedure as the first run was used, except the cells

were infected with cloned MRAd5nef. During both runs, the bioreactors were harvested 48 hours post-infection. Samples were taken and virus concentrations were determined from whole broth (with triton lysis), supernatant, and cell pellets (3 X freeze/thaw) with the AEX and QPA assays. Metabolites were measured with BioProfile 250 throughout the process.

Table 20: Experimental Conditions

Temperature	36.5 ℃	
DO	30%	
PH	7.30	
Agitation	150 rpm	
Sparging	None	<u> </u>

Table 21: Virus source used for experiments.

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Run	Batch ID	Cloned/Uncloned MRKAd5nef	MOI (vp/cells)
#1	B20010115-1	Uncloned	280
	B20010115-2	Uncloned	280
#2	B20010202-1	Cloned	280
	B20010202-2	Cloned	280

Results - Table 22 and 23 show an the ability to scale up production of MRKAd5nef by growth in a bioreactor.

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Table 22: Virus Concentration as measured by the AEX assay

Run	Batch ID	Cloned/Uncloned	Virus Concentration @ 48hpi (1x10 ¹³ vp/L)					
1,011		MRKAd5nef	Supernatant	Clarified Lysate	Total	Triton Lysate		
#1	B20010115-1	Uncloned	0.72	3.26	3.98	5.76		
	B20010115-2	Uncloned	0.38	1.67	2.05	2.46		
#2	B20010202-1	Cloped	0.80	6.00	6.80	8.88		
	B20010202-2	Cloned	0.50	6.00	6.50	8.47		

Table 23: Virus Titers as measured by the QPA assay

Run	Batch ID	Cloned/Uncloned MRKAd5nef	Virus Concentration @ 48hpi (1x10 ¹¹ IU/L)						
,			Whole Broth	Supernatant	Clarified Lysate	Total	Triton Lysate		
#1	B20010115-1	Uncloned	0.13	1.12	1.76	2.88	11.28		
"-	B20010115-2	Uncloned	0.14	0.73	1.54	2.27	5.86		
#2	B20010202-1	Cloned	0.14	0.97	1.62	2.69	11.89		
	B20010202-2	Cloned	0.14	1.17	1.70	2.97	12.47		

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The present invention is not to be limited in scope by the specific embodiments described herein. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art

from the foregoing description. Such modifications are intended to fall within the scope of the appended claims.

EXAMPLE 28

MRKAd5HIV-1gag Boosting of DNA-Primed Animals

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Groups of 3-5 rhesus macaques were immunized with (a) 5 mgs of V1Jns-Flgag (pVIJnsCMV(no intron)-FL-gag-bGHpA), (b) 5 mgs of V1Jns-Flgag formulated with 45 mgs of a non-ionic block copolymer CRL1005, or (c) 5 mgs of V1Jns-Flgag formulated with 7.5 mgs of CRL1005 and 0.6 mM benzalkonium chloride at weeks 0, 4, and 8. All animals received a single dose of 10e7 viral particles (vp) of the MRKAd5HIV-1gag at week 26. Note: 10e7 is too low to prime or boost effectively when used as a single modality (dose is selected to mimic preexposure to adenovirus); see Figure 32.

Blood samples were collected from all animals at several time points and peripheral blood mononuclear cells (PBMCs) were prepared using standard Ficoll method. The PBMCs were counted and analyzed for gamma-interferon secretion using the ELISpot assay (Table 24). For each monkey, the PBMCs were incubated overnight either in the absence (medium) or presence of a pool (called "gag H") of 50 20-aa long peptides that encompass the entire HIV-1 gag sequence.

The results indicate that MRKAd5HIV-1gag was very effective in boosting the T cell immune responses in these monkeys. At week 28 or 2 weeks after the viral boost, the number of gag-specific T cells per million PBMCs increased 2-48 fold compared to the levels observed at week 24 or 2 weeks prior to the boost.

The PBMCs were also analyzed by intracellular gamma-interferon staining prior to (at week 10) and after the MRKAd5gag boost (at week 30). The results for select animals are shown on Figure 31. The results indicate that (a) immunization with DNA/adjuvant formulation elicited T cell responses which can either be balanced, CD4⁺-biased or CD8⁺-biased, and (b) boosting with the MRKAd5gag construct produced in all cases a strongly CD8⁺-biased response. These results suggest that boosting with MRKAd5HIV-1gag construct is able to improve the levels of antigen-specific CD8⁺ T cells.

255 88 88 88 88 25.5 25.5 25.5 25.5 25.5 25.5 828 828 183 828 2-020 **₽₽~~**₹ 959 1915 836 1549 1229 8 5 8 8 5 8 2000 ន % % 22828 85882 និទិនិទ្ធនិ 化妆的 4 0 0 0 0 22 68 88 270 184 154 158 158 158 25 2 2 2 8 8 5 2 2 8 9-00-254 288 119 284 135 149 316 2 49 5 B 원 등 8 85528 **露立っ名**読 400\Z Number of SFChnillion PBMCs

Graf TeO, 4, 8 wts

Table 24, Boost Monks

Table 25, Boost Mon 04020 OCIC OCIK AW3P CB5F AK8B AWZD CAAR CB58 CB5W CB7O MPRCAdSpag(E3+) 10×7 vp MFKAdSgag(E3+) CRL1006/7.5 mgs + 0.6 mM BAK DNA/5mgs + CRL1005/45mgs NA, not available

EXAMPLE 29

Construction of gagpol fusion for MRKAd5gagpol fusion constructs

The open reading frames for the codon-optimized HIV-1 gag gene was fused directly to the open reading frame of the IA pol gene (consisting of RT, RNAseH and integrase domains) by stepwise PCR. Because the gene (SEQ ID NO: 38) does not include the protease gene and the frameshift sequence, it encodes a single polypeptide of the combined size of p55, RT, RNAse H and integrase (1350 amino acids; SEQ ID NO: 39).

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The fragment that extends from the BstEII site within the gag gene to the last non-stop codon was ligated via PCR to a fragment that extends from the start codon of the IApol to a unique BamHI site. This fragment was digested with BstEII and BamHI. Construction of gag-IApol fusion was achieved via three-fragment ligation involving the PstI-BstEII gag digestion fragment, the BstEII/BamHI digested PCR product and long PstI/BamHI V1R-FLpol backbone fragment.

The MRKAd5-gagpol adenovirus vector was constructed using the BglII fragment of the V1R-gagpol containing the entire ORF of gag-IApol fusion gene.

EXAMPLE 30

Immunogenicity Studies in Non-Human Primates

Cohorts of three (3) macaques were immunized with 10e8 or 10e10 viral particles (vp) of one of the following MRKAd5 HIV-1 vaccines: (1) MRKAd5gag; (2) MRKAd5pol; (3) MRKAd5nef; (4) a mixture containing equal amounts of MRKAd5gag, MRKAd5pol, and MRKAd5nef, or (5) a mixture of equal amounts of MRKAd5gagpol and MRKAd5nef. The vaccines were administered at weeks 0 and 4.

The T cell responses against each of the HIV-1 antigens were assayed by IFN-gamma ELISpot assay using pools of 20-aa peptides that encompass the entire protein sequence of each antigen. The results (Table 25) are expressed as the number of spot-forming cells (sfc) per million peripheral blood mononuclear cells (PBMC) that respond to each of the peptide pools.

Results indicate the following observations: (1) each of the single gene constructs (MRKAd5gag, MRKAd5pol, or MRKAd5nef) is able to elicit high levels of antigen-specific T cells in monkeys; (2) the single-gene MRKAd5 constructs can be mixed as a multi-cocktail formulation capable of eliciting very broad T cell responses against gag, pol, and nef; (3) the MRKAd5 vector expressing the fusion

protein of gag plus IA pol is capable of inducing strong T cell responses to both gag and pol.

Table 25. Evaluation of Mixtures of MRKAd5 vectors expressing humanized

HIV-1 gag, pol. gagpol, nef in rhesus macaques

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Grp#	Vaccine	Monk#		T=6 wks				
•	T=0, 4 wks		Mock	Gag H	Pol - 1	Pol-2	Nef	
1	MRKAd5 gag	CB9V	0	15	-	-	-	
	10^10 vp	CD19	0.	374	-	•	-	
	·	109H	1	843	-	•	•	
2	MRKAd5 gag	99D130	1	948	•	-	•	
	10^8 vp	W277	16	324	-	•	-	
		143H	4	595	-	-	•	
3	MRKAd5 pol	CC1X	4		46	256	•	
•	10^10 vp	AW3W	3	-	463	550	•	
		AV43	6	-	95	1333	-	
4	MRKAd5 pol	AW38	1	 	19	30	-	
	10^8 vp	CC8K	0	-	50	995	-	
1		CC21	1] -	33	436	-	
5	MRKAd5 nef	076Q	9	 -	-	·	1204	
	10^10 vp	091Q	4	-	-	•	85	
	·	083Q	0	-			176	
6	MRKAd5 nef	00C029	1	-	-	-	114	
· 1	10^8 vp	98D022	6	\ -	-	-	170	
		98D160	3	-	-	-	198	
7	MRKAd5gag+MRKAd5pol+MRKAd5nef	99D251	3	206	15	193	120	
	10^10 vp each	05H	3	135	21	9	638	
		00C016	3	26	4	51	23	
8	MRKAd5gag+MRKAd5pol+MRKAd5nef	99D215		171	18	193	240	
	10^8 vp each	81H	5	73	6	14	243	
		12H	8	1140	115	811	719	
9	MRKAd5gagpol +MRKAd5 nef	99D211	0	83	56	838	725	
	10^10 vp each	22H	4	385	119	1194	1915	
		61H	4	343	11	765	853	
10	MRKAd5gagpol +MRKAd5 nef	34H	3	78	19	5	75	
l	10 ^A 8 vp each	48H	1	65	105	46	43	
	•	70H	5	158	15	220	191	

Indicated are numbers of spot-forming cells per million PBMCS against the peptide pools. Mock, no peptides; gag H, fifty 20-aa peptides encompassing p55 sequence; pol-1, 20-aa peptides representing N-terminal half of IA pol; pol-2, 20-aa peptides representing the carboxy-terminal half of IA pol; nef, 20-aa peptides encompassing the entire wild-type nef sequence. Responses to the antigens prior to the first immunization did not exceed 40 sfc/10⁶ PBMC.

WHAT IS CLAIMED IS

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A recombinant adenoviral vaccine vector at least partially deleted in
 and devoid of E1 activity, comprising:

- a) an adenovirus cis-acting packaging region corresponding to from about base pair 1 to between from about base pair 400 to about base pair 458 of a wildtype adenovirus genome; and
- b) a gene encoding an HIV protein or immunologically relevant modification thereof.
- 2. A vector in accordance with claim 1 comprising a packaging region corresponding to from about base pair 1 to about base pair 450 of a wildtype adenovirus genome.
- 3. A vector in accordance with claim 1 further comprising nucleotides
 15 corresponding to between from about base pair 3511 to about 3524 to about base pair
 5798 of a wildtype adenovirus genome.
 - 4. A vector in accordance with claim 3 comprising base pairs corresponding to 1-450 and 3511-5798 of a wildtype adenovirus genome.
- 5. A vector in accordance with claim 4 which is deleted of base pairs20 451-3510.
 - 6. A vector in accordance with claim 1 which is at least partially deleted in E3.
 - 7. A vector in accordance with claim 6 wherein the E3 deleted region is from base pairs 28,133-30,818.

8. A vector in accordance with claim 1 wherein the gene encoding the HIV protein or modification thereof comprises codons optimized for expression in a human.

- 9. A vector in accordance with claim 1 wherein the vector comprises a gene expression cassette comprising:
 - a) a nucleic acid encoding a protein;

- b) a heterologous promoter operatively linked to the nucleic acid encoding the protein; and
 - (c) a transcription termination sequence.
- 10. A vector in accordance with claim 9 wherein the gene expression cassette is inserted into the E1 region.
 - 11. An adenoviral vector in accordance with claim 9 wherein the gene expression cassette is in an E1 parallel orientation
- 12. An adenoviral vector in accordance with claim 9 wherein the geneexpression cassette is in an E1 antiparallel orientation.
 - 13. An adenoviral vector in accordance with claim 9 wherein the promoter is a cytomegalovirus promoter devoid of intronic sequences.
 - 14. An adenoviral vector in accordance with claim 13 wherein the promoter is an immediate early human cytomegalovirus promoter.
- 20 15. An adenoviral vector in accordance with claim 9 wherein the promoter is a murine cytomegalovirus promoter.
 - 16. An adenoviral vector in accordance with claim 9 wherein the transcription termination sequence is a bovine growth hormone polyadenylation and transcription termination sequence.

17. An adenoviral vector in accordance with claim 9 wherein the transcription termination sequence is a synthetic polyadenylation signal (SPA).

- 18. A cell comprising the adenoviral vector of claim 1.
- 19. Recombinant, replication-defective adenovirus particles harvested
 and purified subsequent to transfection of the adenoviral vector of claim 1 into a cell
 line which expresses adenovirus E1 protein at complementing levels.
 - 20. An HTV vaccine composition comprising purified adenovirus particles of claim 19.
- 21. An HIV vaccine composition of claim 20 which comprises aphysiologically acceptable carrier.
 - 22. A method of producing recombinant, replication defective adenovirus particles containing the adenoviral genome of the adenoviral vector of claim 1 which comprises introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and harvesting the resultant recombinant, replication-defective adenovirus.
 - 23. A method according to claim 22 wherein the cell is a PER.C6[®] cell.

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- 24. A method of generating a cellular-mediated immune response against HIV in an individual comprising administering to the individual a vaccine of claim 21.
- 25. A method according to claim 24 which further comprises administration to the individual a DNA plasmid vaccine, optionally administered with a biologically effective adjuvant, protein or other agent capable of increasing the immune response.

26. A method according to claim 25 wherein the DNA plasmid vaccine is administered to the individual prior to administration of an adenovirus vaccine.

- 27. A method according to claim 24 wherein the adenovirus vaccine is
 5 preceded by an adenovirus vaccine of a different serotype.
 - 28. A method according to claim 24 which comprises administering and readministering the adenovirus vaccine vector to the individual.
 - 29. An adenoviral vector in accordance with claim 1 wherein the HIV protein is HIV gag or an immunologically relevant modification thereof.
- 30. An adenoviral vector in accordance with claim 9 wherein the gene expression cassette comprises an open reading frame encoding an HIV gag protein or immunologically relevant modification thereof.
 - 31. A recombinant adenoviral vaccine vector at least partially deleted in E1 and devoid of E1 activity, comprising:
- a) an adenovirus *cis*-acting packaging region corresponding to from about base pair 1 to about base pair 450 and from about 3511 to about 5798 of a wildtype adenovirus genome, and deleted for base pairs corresponding to from about base pair 451 to from about base pair 3510 of a wildtype adenovirus genome; and
 - b) a gene expression cassette comprising
 - i) SEQ ID NO: 29;
 - ii) a heterologous promoter operatively linked to i); and
 - iii) a transcription termination sequence.

32. An adenoviral vector in accordance with claim 31 wherein the gene expression cassette is in an E1 parallel orientation.

- 33 An adenoviral vector in accordance with claim 31 wherein the gene expression cassette is in an E1 antiparallel orientation.
- 34. An adenoviral vector in accordance with claim 31 wherein the promoter is a cytomegalovirus promoter devoid of intronic sequences.

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- 35. An adenoviral vector in accordance with claim 31 wherein the transcription termination sequence is a bovine growth hormone polyadenylation and transcription termination sequence.
- 36. An adenoviral vector in accordance with claim 31 which is at least partially deleted in E3.
 - 37. A cell comprising the adenoviral vector of claim 30.
 - 38. Recombinant, replication-defective adenovirus particles harvested and purified subsequent to transfection of the adenoviral vector of claim 30 into a cell line which expresses adenovirus E1 protein at complementing levels.
 - 39. An HIV vaccine composition comprising purified adenovirus particles of claim 38.
 - 40. An HIV vaccine composition of claim 39 which comprises a physiologically acceptable carrier.
 - 41. A method of producing recombinant, replication defective adenovirus particles containing the adenoviral genome of the adenoviral vector of claim 30 which comprises introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and harvesting the resultant recombinant, replication-defective adenovirus.

42. A method according to claim 41 wherein the cell is a PER.C6[®] cell.

43. A method of generating a cellular-mediated immune response against HIV in an individual comprising administering to the individual a vaccine of claim 21.

- 44. A method according to claim 43 which further comprises administration to the individual a DNA plasmid vaccine, optionally administered with a biologically effective adjuvant, protein or other agent capable of increasing the immune response.
- 45. A method according to claim 44 wherein the DNA plasmid vaccine is administered to the individual prior to administration of an adenovirus vaccine.
 - 46. A method according to claim 43 wherein the adenovirus vaccine is preceded by an adenovirus vaccine of a different serotype.
- 47. A method according to claim 43 which comprises administering and readministering the adenovirus vaccine vector to the individual.
 - 48. An adenoviral vector in accordance with claim 1 wherein the HIV protein is HIV pol or an immunologically relevant modification thereof.
- 49. An adenoviral vector in accordance with claim 9 wherein the gene
 20 expression cassette comprises an open reading frame encoding an HIV pol protein or immunologically relevant modification thereof.
 - 50. A recombinant adenoviral vaccine vector at least partially deleted in E1 and devoid of E1 activity, comprising:

a) an adenovirus cis-acting packaging region corresponding to from about base pair 1 to about base pair 450 and from about 3511 to about 5798 of a wildtype adenovirus genome, and deleted for base pairs corresponding to from about base pair 451 to from about base pair 3510 of a wildtype adenovirus genome; and

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- b) a gene expression cassette comprising
 - i) a nucleotide sequence selected the group consisting of SEQ ID NO: 1, SEQ ID NO: 5 and SEQ ID NO: 7;
 - ii) a heterologous promoter operatively linked to i); and
 - iii) a transcription termination sequence.
- 51. An adenoviral vector in accordance with claim 50 wherein the gene expression cassette is in an E1 parallel orientation.
- 52. An adenoviral vector in accordance with claim 50 wherein the gene expression cassette is in an E1 antiparallel orientation.
- 53. An adenoviral vector in accordance with claim 50 wherein the promoter is a cytomegalovirus promoter devoid of intronic sequences.
- 54. An adenoviral vector in accordance with claim 50 wherein the transcription termination sequence is a bovine growth hormone polyadenylation and transcription termination sequence.
- 55. An adenoviral vector in accordance with claim 50 which is at least partially deleted in E3.
 - 56. A cell comprising the adenoviral vector of claim 49.

57. Recombinant, replication-defective adenovirus particles harvested and purified subsequent to transfection of the adenoviral vector of claim 49 into a cell line which expresses adenovirus E1 protein at complementing levels.

- 58. An HIV vaccine composition comprising purified adenovirus5 particles of claim 57.
 - 59. An HIV vaccine composition of claim 58 which comprises a physiologically acceptable carrier.
 - 60. A method of producing recombinant, replication defective adenovirus particles containing the adenoviral genome of the adenoviral vector of claim 49 which comprises introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and harvesting the resultant recombinant, replication-defective adenovirus.

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- 61. A method according to claim 60 wherein the cell is a PER.C6® cell.
- 62. A method of generating a cellular-mediated immune response against HIV in an individual comprising administering to the individual a vaccine of claim 59.
 - 63. A method according to claim 62 which further comprises administration to the individual a DNA plasmid vaccine, optionally administered with a biologically effective adjuvant, protein or other agent capable of increasing the immune response.
 - 64. A method according to claim 63 wherein the DNA plasmid vaccine is administered to the individual prior to administration of an adenovirus vaccine.

65. A method according to claim 62 wherein the adenovirus vaccine is preceded by an adenovirus vaccine of a different serotype.

- 66. A method according to claim 62 which comprises administering and readministering the adenovirus vaccine vector to the individual.
- 67. An adenoviral vector in accordance with claim 1 wherein the HIV protein is HIV nef or an immunologically relevant modification thereof.

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- 68. An adenoviral vector in accordance with claim 9 wherein the gene expression cassette comprises an open reading frame encoding an HIV nef protein or immunologically relevant modification thereof.
- 69. A recombinant adenoviral vaccine vector at least partially deleted in E1 and devoid of E1 activity, comprising:
 - a) an adenovirus *cis*-acting packaging region corresponding to from about base pair 1 to about base pair 450 and from about 3511 to about 5798 of a wildtype adenovirus genome, and deleted for base pairs corresponding to from about base pair 451 to from about base pair 3510 of a wildtype adenovirus genome; and
 - b) a gene expression cassette comprising
 - i) a nucleotide sequence selected the group consisting of SEQ ID NO: 9, SEQ ID NO: 11, SEQ ID NO: 13 and SEQ ID NO: 15;
 - ii) a heterologous promoter operatively linked to i); and
 - iii) a transcription termination sequence.
- 70. An adenoviral vector in accordance with claim 69 wherein the gene expression cassette is in an E1 parallel orientation.

71. An adenoviral vector in accordance with claim 69 wherein the gene expression cassette is in an E1 antiparallel orientation.

- 72. An adenoviral vector in accordance with claim 69 wherein the promoter is a cytomegalovirus promoter devoid of intronic sequences.
- 73. An adenoviral vector in accordance with claim 69 wherein the transcription termination sequence is a bovine growth hormone polyadenylation and transcription termination sequence.

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- 74. An adenoviral vector in accordance with claim 69 which is at least partially deleted in E3.
 - 75. A cell comprising the adenoviral vector of claim 68.
- 76. Recombinant, replication-defective adenovirus particles harvested and purified subsequent to transfection of the adenoviral vector of claim 68 into a cell line which expresses adenovirus E1 protein at complementing levels.
- 77. An HIV vaccine composition comprising purified adenovirus particles of claim 76.
 - 78. An HIV vaccine composition of claim 77 which comprises a physiologically acceptable carrier.
 - 79. A method of producing recombinant, replication defective adenovirus particles containing the adenoviral genome of the adenoviral vector of claim 68 which comprises introducing the adenoviral vector into a host cell which expresses adenoviral E1 protein, and harvesting the resultant recombinant, replication-defective adenovirus.
 - 80. A method according to claim 79 wherein the cell is a PER.C6® cell.

81. A method of generating a cellular-mediated immune response against HIV in an individual comprising administering to the individual a vaccine of claim 78.

- 82. A method according to claim 81 which further comprises

 5 administration to the individual a DNA plasmid vaccine, optionally administered with
 a biologically effective adjuvant, protein or other agent capable of increasing the
 immune response.
- 83. A method according to claim 82 wherein the DNA plasmid
 vaccine is administered to the individual prior to administration of an adenovirus
 vaccine.
 - 84. A method according to claim 81 wherein the adenovirus vaccine is preceded by an adenovirus vaccine of a different serotype.
 - 85. A method according to claim 81 which comprises administering and readministering the adenovirus vaccine vector to the individual.
- recombinant, replication-defective adenovirus particles, wherein the adenovirus particles are harvested and purified from a cell line expressing adenovirus E1 protein, and wherein the particles are harvested subsequent to transfection of the cells with an adenoviral vector or vectors in accordance with claim 9; said vector(s) comprising a gene expression cassette or cassettes comprising nucleotide sequences encoding HIV proteins selected from the group consisting of:
 - a) gag, pol, and nef, expressed independently from three individual vectors;

 b) gag, pol, and nef, expressed independently from one vector with the encoding nucleic acid sequences operatively linked to distinct promoters and transcription termination sequences;

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 gag, pol, and nef, expressed via two vectors, one expressing a polnef fusion, and another expressing gag;

- d) gag, pol, and nef, expressed via two vectors, one expressing a gagpol fusion and another expressing nef;
- e) gag, pol and nef, expressed via two vectors, one expressing a nefgag fusion and another expressing pol;

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- f) gag, pol, and nef, expressed via one vector expressing a gag-polnef fusion;
- g) gag and pol, expressed independently from two individual vectors;
- h) gag and pol, expressed independently from one vector with the encoding nucleic acid sequences operatively linked to distinct promoters and transcription termination sequences;

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- i) pol and nef, expressed independently from two individual vectors;
- j) pol and nef, expressed independently from one vector with the encoding nucleic acid sequences operatively linked to distinct promoters and transcription termination sequences;

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- k) nef and gag, expressed independently from two individual vectors;
- nef and gag, expressed independently from one vector with the encoding nucleic acid sequences operatively linked to distinct promoters and transcription termination sequences;
- m) gag and pol, expressed via one vector expressing a gag-pol fusion;

n) pol and nef, expressed via one vector expressing a pol-nef fusion; and

- o) nef and gag, expressed via one vector expressing a nef-gag fusion.
- 87. A multivalent adenovirus vaccine composition in accordance with claim 86 wherein the gag-pol fusion consists of SEQ ID NO: 39.
 - 88. A multivalent adenovirus vaccine composition in accordance with claim 86 wherein the fused sequences have the encoding nucleic acid sequences operatively linked to distinct promoters and transcription termination sequences.
- 89. A multivalent adenovirus vaccine composition in accordance with

 10 claim 86 wherein the fused sequences have the encoding nucleic acid sequences

 operatively linked to a single promoter; and the encoding nucleic acid sequences

 operatively linked by an internal ribosome entry sequence ("IRES").

Original Adenovector Construct:

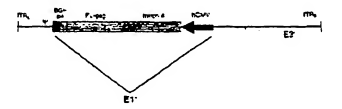


Figure 1: Original HIV-1 gag adenovector.

Sequence of the open reading frame for FL-gag (human codon optimized)

atgggtgctagggcttctgtgctgtctggtggtgagctggacaagtgggagaagatcaggctgaggcctggtgg caagaagaagtacaagctaaagcacattgtgtgggcctccagggagctggagaggtttgctgtgaaccctggc agctgaggtccctgtacaacacagtggctaccctgtactgtgtgcaccagaagattgatgtgaaggacaccaag gaggccctggagaagattgaggaggagcagaacaagtccaagaagaaggcccagcaggctgctgctggc acaggcaactccagccaggigtcccagaactaccccattgigcagaacctccagggccagatggtgcaccag gecatetececeggaccetgaatgeetgggtgaaggtggaggagaaggeetteteeetgaggtgateee catottctctqccctgtctgagggtgccacccccaggacctgaacaccatgctgaacacagtgggggggccatc aggetgecatgeagatgetgaaggagaecateaatgaggaggetgetgagtgggaeaggetgeateetgtge acgciggccccattgcccccggccagaigagggagcccaggggctctgacattgctggcaccacctccaccct ccaggagcagattggctggatgaccaaccaccccccatccctgtgggggaaatctacaagaggtggatcat ccttcaqqqactatgtqgacaggttctacaagacctgagggctgagcaggctcccaggaggtgaagaact ggatgacagagaccctgctggtgcagaatgccaaccctgactgcaagaccatcctgaaggccctgggccctg ctoccaccttggaggagatgatgacagcttgccagggggtggggggccttggtcacaaggccagggtgctg gctgaggccatgtcccaggtgaccaactccgccaccatcatgatgcagaggggcaacttcaggaaccagag gaagacagtgaagtgcttcaactgtggcaaggtgggccacattgccaagaactgtagggcccccaggaaga ggcaaaatctggccctccacaagggcaggcctggcaacttcctccagtccaggcctgagcccacagccct ageigiaeceeciggeeicecigaggieecigttiggeaacgaeceelecieecaglaaaalaaageecgggea gat (SEQ ID NO: 29)

Figure 2

Old Transgene: New Transgenes: New GAG BOH NCMV GAG SPA

Figure 3: Diagrammatic representation of the original HIV-1 gag transgene and the series of new transgene constructions.

GAG

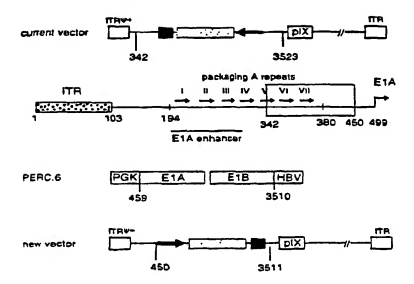


Figure 4: Modifications made to the current adenovector backbone in the generation of the new vector.

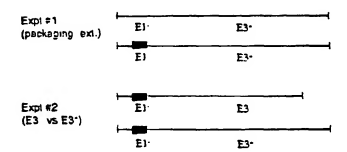


Figure 5: Virus mixing experiments to determine the effects of the addition made to the packaging signal region (Expt #1) and analysis of the effects of the E3 gene on viral growth (Expt. #2). The red bars denote the region of modifications made to the E1 deletion.



Figure 6: Autoradiograph of viral DNA analysis following viral mixing experiments (expts. #1 and #2) as detailed in the text.

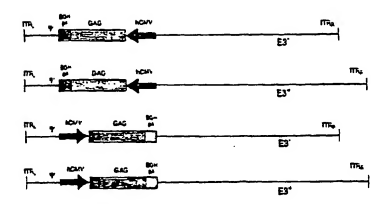


Figure 7A: hCMV-FLgag-bGHpA adenovectors constructed within the *MRK* backbone. E1 parallel and E1 antiparallel transgene orientation within the E3- and E3+ backbones were constructed.

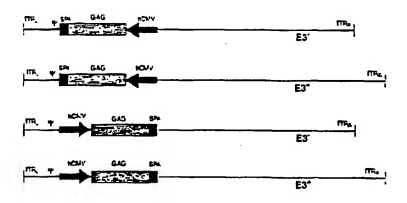


Figure 7B: hCMV-FLgag-SPA adenovectors constructed within the "MRK" backbone. E1 parallel and E1 antiparallel transgene orientation within the E3- and E3+ backbones were constructed.

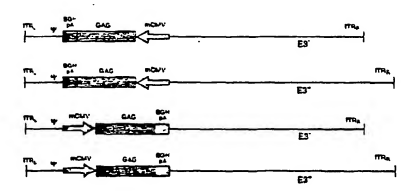


Figure 7C: mCMV-FLgag-bGHpA adenovectors constructed within the *MRK* backbone. E1 parallel and E1 antiparallel transgene orientation within the E3- and E3+ backbones were constructed.

Plasmid mixing expt: (orientation)

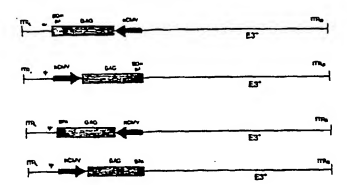


Figure 8A: Effect of transgene orientation

Plasmid Mixing expt: (poly A signal)

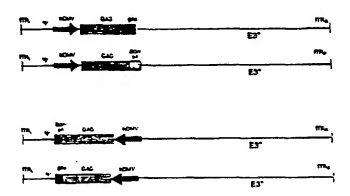


Figure 8B: Effect of polyadenylation signal



Figure 9: Viral DNA from the four Adgag candidates at P5, following BstE11 digestion.

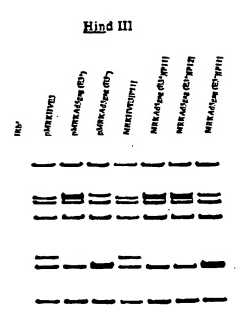


Figure 10: Viral DNA analysis of passage 11 and/or 12 of MRKHVE3, MRKAd5gag and MRKAd5gag(E3-).

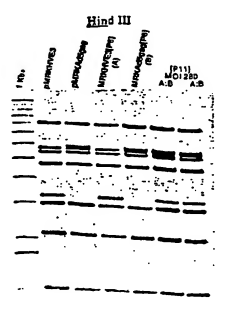


Figure 11: Viral DNA analysis (*Hin*dIII digestion) of passage 6 MRKHVE3 and MRKAd5gag used to initiate the viral competition study. Last two lanes are passage 11 analysis of duplicate passages of the competition study (each virus at MOI 280 vp).

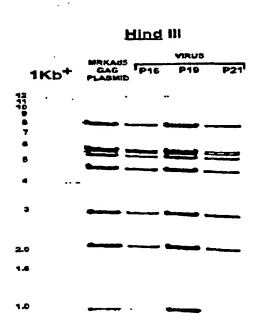
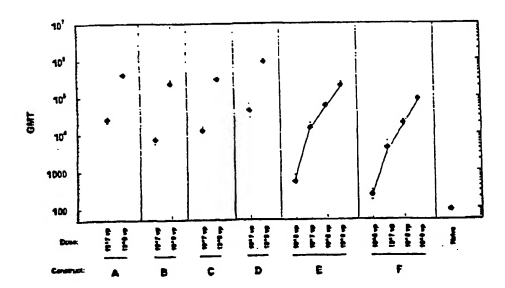


Figure 12: Viral DNA analysis by *Hind*III digestion on high passage numbers for MRKAd5gag in serum containing media with collections made at specified times. The first lane shows the 1 Kb DNA size marker. The other lanes represent pre-plasmid control (digested with Pac1 and *Hind*III), and MRKAd5gag virus continually passaged to P16, P19 and P21(serum containing media).

Figure . Serum anti-p24 Levels at 3 Wks post i.m. immunization of balb'c mice (n=10) with Varying Doses of Several Adgag constructs: (A) MRKAd5gag (through passage 5): (B) MRKAd5 E3' hCMV-FLgag-bGHpA; (C) MRKAd5 E3' hCMV-FLgag-SPA; (D) MRKAd5 E3' mCMV-FLgag-bGHpA; (D) research Lot (293 cell-derived) of Ad5HIV-lgag; and (F) clinical lot (Ad5gagFN0001) of Ad5HIV-lgag. Reponed are the geometric mean titers (GMT) for each cohort.



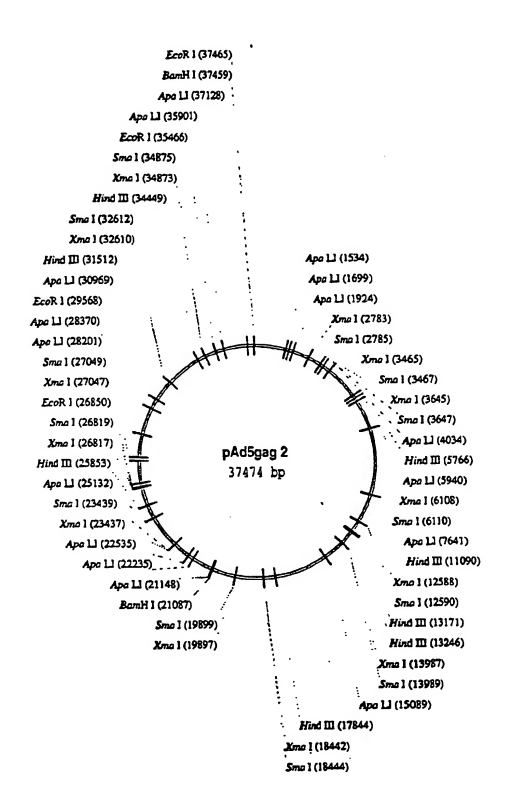


Figure 14

כוכופטכנייד CCTATAGTA CATTAGETICA GTANTCANGT AATCACCTAT TINCTOCATA AGTACATOTA TCATGOTAGAT CCATCCGCAC CACTATAGAGAGG CHECACTERIA CCATATCATY ATAGTATAC CCACCACTIC CACCTTACITY TATCATATY **GCTANAYCOT** TOGAGGICCC GOTCTACCAC TOTOCACC ACATCAAGTO AGGCCTTCTG COGTANAAGC COTTTACCTO TACGETICITAT ATCCAACATA ATTACGGGGT TAATGCCCCA TOTACTTCAC GGATGAACCG CATTICCAND CTANAGGETIC COTITACCCO CTCCATAGAA GOAGGTCCCT AGGCTCTGAG ACCTCCAGOO ACTOCANANA OCCARTITICO GCANATGCAC TOACGTCAAT ACTOCAGITA CCTACTTOGC GAGGTATCTT TCCCCAAGAC TCCGAGACTC ATTOROGOGO TGACGITTIT **GCANATEGGG** CCTCCAGGGA CATTICTARAC GGACTITIGAC CCTCAMCTO ATCCATTICCA GROCCOCOTA ATCCCACTIT COCOTANCTO CTUTTTTOAC CACAAAACTO TRANSATICTAE CATGGGTGCT GTACCCACGA ATTICTUTES TANCACACCC CCCTCCANAC COGACOTITO CCTOGAGAAG CONTROLLEG GGACCICITIC ATTOTOTOCAGA TAACACGTCT TACCTACCT ATAGTAATCA TATICATTAGE CCCCCCCCAT TOMOGOSTICA TACCETGANA GACTICACOOG CTCAGTGCCC CCCCATTGAC CTCCCANANG CACCOTITIC GTAAGATTTG ACTTOCCAOT THOTOTOT AAACACTGCA GANCTACCCC TATCCCCANA ANCANCTECE CCTAAAGCAC GAGCTCGGGA CTACTTATTA ACATGACCTT ATAGGGGGTTT GREAGTETAG COGNICITETS COSTARSTEC ACTETAGATO CGATTICGTG CCTATACCAG AACACATGTA ACKIGACGAT TCACTCCCTA CARGITAACCGA CCATATTICATE אסגיניכניניספ TCCCGGCGCC CHICKLETT TATTATTATA GREGEEGE GATCAATAAT GCCCAACGAC CGGGTTGCTG TAMACTICCCC ATTITISACTIO TOTACTOOM THEFTIGAGE GCCATCCACG CTCCAGCCCT CTANGGAC CTTCATCOCC נירניכאניכדכ 2020022522 CANCERCAM ATANTANTAT GCCTGGAGAC TACHGRACAT GTAMITTIGG TATKY ICCCAT TATAAACAGA PCATTATICA CACCCIACTOR ATACCCCTICA TACCCACACC AAAATISTOGT CHINCHARIAG CACCASTICTO MCANCTACAA TYTHICATES דאיאמידית איכאנאפעיד CATTFAAACC ATATTTGTCT ACTAATAACT CHARITAGE CITATITACIA CATANATIAL TATGCCCAGT ATIGGRACITIES TTTTACAGCA CCTGGGGCCAG CATACCCCACTIC CTTCANTIACA COLOR CACTRICATOR ACTIVITICOCA ATCATAATCA TACTATTACT TTACCGGGG CASTACATEA CCCTGAAACA CCAGGCAGAT AACTICACITY TACAACTOTA ANTINGCUCC CCCGACCCTA GTCATUTAGE ניבטוביאטעוב COGATITICCCC GCCTAACACG COTTOTATA CCACCGTTCT CONCOUNTER CANCATTUAT CTINTANTA GCATGITAGIA ATCHINACAT ANTORGENISA CCCCTTCCCAT CAGGACTITICC TANCTICANT ACTIV YASTI'A ACT TO THE POSC 16 TCACACCGCC CCTACAACAT ATAXXXCGTA TTACCCACCT ACCGRETCE CATTACCGCC ACTTACGGTA OFMCTGCAG CCCCTTTTCC CAAAACCGTG GTTTTAGTTG CCADATATAT TEGTETEGAG CANATENETT CCACTCCCCA **TETCAGGGGT** TATTTAGAT ATAAAACCTA TO TOTAL COTTTTACC CCANANTCCG PUTCHTACTC ACACAATGAG CCOGGTCAM GCCCAGTTT GTAATGGCGG TCAATCCCAT CATTRIACRITE CTAAAATCCC CATTTACCOS OCTACCACTA COCCANANCC CAMATCANC GTTTAGTGAA CATTOONACG CCCTTGCCAC GTAACCTTGC **OCTONOCICE** AGACTCCCCA CHCTVXCACCA CACACCTROFT TOTACACIC ACTACAACGT TECETIONNO ACCAGAGETE AGAAGATCAG TOGOACATGA MACRICICIA **GTCAATGACG** CCATCCTCAT GOGNACOOTO TCTTCTAGTC **GCTOGAGACC** ACCENCANCE APARCETAC TCCGACGACG **GCCTTACATA** CHITTOGCAC COACCTCTOO TAMARGORIS ACTTATTAM TCATCTCCAA COCNATIONAL ACCONCITITO CAGTTACTOC ATTATATATA ההכסואהחה CCCCCTTCAC ATTITION TCAATAATT THECOGRAFT AGTACAGGIT TAATATACT TACCCTCAAA **OCTETATATA GTOTCACCOA** MODECCAGE Treconstea AGAGTCCACA ATATACCO TACCICANGO TAACGCCAAT ATTOCOGITA CCCTATTGAC ATCCCTATTA TACCGATAAT ATCOCACTTT בעככסכסככ GACCCCCCC GACAAGTGGG TOCOACCOOR **NACTONNATC** ATCOACTICC GOCATAACTO CHOTTICACCC ACCETOOCCT CACAGTOOCT CATCATCACA COAAGTGACA CCTTCACTGT TICACTITIAG TTATATTGGC TTCTTAATTA ACATCATCAA TRETACTAGET GTAGTAGTGT Tercaggigt ATGCCACCCT GANTARGAGG **TATACATOTA** VTCGGGGTATA GITCCCATAG CCCTOTACAA CAGGITCTIC CACCOCCICAC CCCCCACTG DOTOTACACA CCACATGTOF CTTATTCTCC GTCCACAAAA PAGCCCATAT CAAGGGTATC CANDINCOCC STICATOCOO COTATTAGT **SCATAATCAG** NTCACCICA **PACTOCADT** PACCOTOCOGA CCGATCCAGC DOCTADOTEG NOTCAGCTG ACCACTCOAC TTOCTOTOA MACCIACACT **JOGACATOTT** DICCARRANG ATATGTACAT ANCIANTINAT CAGGIGITIT 1401 1501 601 701 901 1001 1101 1201 1301 1601 501 **B**01 101 201 301 601

Figure ISA

PMRKAdhaaa HER682

1701	CACCAGGCCA	Telececec	CACCCTCAAT	CCCTCTCTCACTCAC	Aritharina;A	נאנאטאאנטניכ	MONGOGONE	MCCACTAGGG	GTACAGGAGA	GCCATACAGAC
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	TCCCACGGTG	GOGGOTICCTO	GACTIVERIAFIE	ACCIAL TITITO	TCNTTYCO.G	CITACITY:CGAC	RCTACGTUTA	CGACTICCTC	TOGTACTTAC	Techniciae
1901	TGACTGGGAC	ACCCTCCATC	CTRITACACK	TRACICCATT	שמעמנונאנוני	אייאייאניאראראייא	CKCCAGGACAC	TCTGACATIG	CTORCACCAC	CTCCACCCI
	ACTCACCCTG	TCCOACGTAG	GACACTETICAL	ACCCARACTAA	נישישרישישישי	ナンストナイントナ	כניים כניכני כני	AGACTGTAAC	GACCOACGIO	かんろうない
2001	CAGGAGCAGA	TROCCTODAT	GACTINATING	COCCCCATC	VIXXIOXXIIV	ANTITACAAG	NYTHANATEA	recreasee.	GANCAAGATT	เสนาสาราสารา
	Greencoter	ANCCOACCTA	CTCGTTCTTG	CONTRACTAGE	האנאניכנית	TTAGATGTTC	TCCACCTAGE	AGGACCCGGA	CTICTION	CACTCCTAIN
2101	ACPCCCCCCAC	CTCCATCCTO	GACATEMARIC	ACXXXTCCAA	משננינות	ACCONCTATO	TGGACAGGTT	CTACAAGACC	CTGAGGGCTG	AGCAROCCT
	TOAGGGGGTG	GAGGTAGGAC	CTGTACTCCG	TCCCGGGGTT	ככונטטאנאענ	TCCCTVATAC	ACCTGTCCAA	GATGTTCTGG	GACTCCCGAC	TCUTCOGA
2201	CCAGGAGGTG	AAGAACTOGA	TCACACACAC	CCTGCTCGTG	CAGAATGCCA	ACCENTACTG	CANGACCATC	CTGAAGGCCC	TOGRECETOC	TOCCACCCT
	OCTECTECAC	TICTICACCT	ACTORCICIO	GGACGACCAC	CICTIACCCT	TOCHACTERE	GITTCTGGTAG	GACTTCCGGG	ACCCCCCCCACC	ACCOURCEON
2301	DAGGAGATCA	TOACACCCTG	CCACCACCATE	SCANSOCICE TO	CTCACAARCC	CARRETTO	GCTGAGACCA	TOPCCCAGGT	DACCAACTEC	מככאונכאובי
	CICCICINC	ACTOTOGGAC	GGTCCCCCAC	CCCCCCCCCC	CACTUTTOCO	GTCCCACCAC	CGACTCCGGT	ACAGGGTCCA	CTCOTTCAGG	COGNOCTAC
2401	TGATGCAGAG	GOCCAACTIC	ACCIDANCEAGA	GRANGACAGT	מאתומנידוכ	AACTIGHTACEA	AGGTGGGCCA	CATTGCCANG	MCTGTAGGG	כככניניאשמא
	ACTACOTOTIC	CCCOTTGAAG	recenteerer	CCTTCTGTCA	CTTCACGAAG	THINCACCUT	TCCACCCAGT	GTAACGGTTC	TTGACATCCC	GOOGGOTCCT
2501	GAAGGOCTOC	TOGAAGTGTG	GCANGGANGG	CCACCAGATO	ANCHACTOCA	ATGAGAGGCA	ACCENACITIC	CTCCCCAAAA	retroceeere	CCNCNACION
	CTIVECCGACG	*	_	REPOSTETAC	TRUCKARGE	TACTCTCCGT	CCGGTTGAAG	OACCCONTIT	AGACCOGGAG	GCTCTTCCCV
2601	AGGCTGGCA	ACTICCTOCA	GTCCAGGCC7	GAGCCCACAG	CCCCTCCCGA	CONDICCTIC	AGGETTICORG	ACCACARCAC	CACCCCAGC	CACIANCICAL
	TCCGGACCGT	F	CAGGTCCGGA	CTCCCCTGTC	GCXXCAGGGCT	CCTCACGOAGG	TCCAACCCC	recterreno	gradocated	FICHTOGICA
									-	<u> </u>
2701	ACTICATION	CAAGGGACTO	TACCCCCTTIO	CCTCCCTGAG	GICCCIGITIF	CHICAARTIACC	CCTCCTCCCA	GTANNITAAA	ОСССОВВЕСТО	ATCHOCTOTA
	TCGGGTAACT		ATOCOCOCACC	האוממשכדכ	CAGGGGACAAA	CCGITTERTEG	COARCARORT	CATITITATIT	coocceete	TAGACGACA
1000		6	A CHARLETTE		TRECTTECT	GACCTOCA	CIGHICCACTC	CCACTOTCCT	FTCCTANTA	ANTERCOAN
1007	CONTRACTOR OF THE REAL		BLANCANACG	GCCAGGGGGC	ACCEAAGGAA	CTGGGACCTT	CCACGGTGAG	GGTGACAGGA	AAGGATTATT	TTACTOCTO
•	50140505	2000								Sphi
1000	Print A Bride	The street of the street	ACROHOTICATE	CTATICTURE	SUSSISSISSISSISSISSISSISSISSISSISSISSISS	CASCACTOACA	CCAACOTATA	CCATTOOGAA	GACAATAGCA	GCX:ATTXCTKX
1009			TCCACAGTAA		CCCACCCCAC	CCCGTCCTGT	COTTCCCCCT	CCTANCCCTT	CTCTTATCCT	CCCTACGACC
			Pwd							
	•		2	P						
3001	GTATOCOOTO	OCCUCATOO	CCCATCGGUG	COCCUTACIO	AAATGTGTT	CATCHOOC TAN	AGGGTGGGAA	ACNATATATA	Acordococr	CITATOTAGE
	CCTACGCCAC	Ç	GOCTAGCCGC	OCCUCATGAC	TTTACACACC	COCACCGAAT	TCCCACCCTT	TCTTATATAT	TCCACCCCCA	CANTACA TEN
									manual and a second	
1101	THETATERE	TTTTGCAGCA	פככפכנעכנפ	CCATGMGCAC	CAACTECTET	GATGGAAGCA	THISTOPHICAC	ATAPPEACA	ACCIOCATOC	CCCCATAGGC
	AACATAGAC	5	כסכיטטכנענ	CETACTICATIC	CTTCACCAAA	CTACCTTACT	AACACTCGAG	TATABACTOF	TUCCICOTACO	COCCTACCO!
1000	The state of the s	Ľ	Transact Transact	CATRIARY	ددردردردر	TERCETATANA	בבבדאינדאינ	TTCACCTACG	AGACCCTOTC	TOCARCCCCG
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figure 158

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Figure 150

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1064	OCTOCOCTTO TOTAL STRANG	AGOCTOGEC	TACTOSTICET ACCIDENCACEA	GAARAXATKIC	CONTRACTOR	נינדנאנאביה היאבנאנאנאנאפ	CCCACTING	CATTTCACCA	TOCHOTCATA ACCACAGINT	GYCCAGGGGG CAGGTCGGGG
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	AGGCGCCCCA	CCGGGAACCG	CCCOTCGAAC	CCCONCCTCC	Treamant	נא ידי,כתיכפדכ	ACCITCIANA	ACTCCCCCAT	CTCGAACCCG	CCCTCTTTAT
5101	CCGATTCCGG	CETCATECOT	TCCACGCCGC Aggccgcgcg	AGGCCCCACA	CTRCCAGAGG	CATTCCACGA	COCHECACITE	CHCTGGCCGT	TCTGGGGTCAA AGCCCCAGTT	AAARTCAGGTT TITTGGTCCCA!
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5301	AGAGGCCTOT	•	TOTACCIACOS	TECTOCTOR	ATAGMACTC	GACCACTCT	CACACAAAGG	CICCCCATCCA	DOCCAGCACO	AACKINCICTA
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5401	ACTOCCADCO	OTACCOCICO	TRITCCACTA	COCCUTECAE	TURCHECAGG	MIGHGAAGAC	ACATGTCGCC	CICITOGGCA	TCANGGAAGO	TOATTOSTET
	TCACCCTCCC	CATCGCCAGC	AACAGGTGAT	CCCCCAGGTU	ACCOMPUTED	CACACTTCTO	TOTACAGOOG	GAGAAGCCGF	AGTICCITICC	ACTANCCANA
5501	CTAGGTGTAG	OCCACOTOAC	COGGRANTICC	TCAACCCCCCC	CTATAAAACG	GACTICAGGGC	acerrearce	TCACTCTCT	CCOCATCOCT	GTCTGCGAGG
	CATCCACATC		GCCCACAAGG	ACTICCCCCC	GATATITICS	CCCACCCTCG	CCCAARCAGG	ACTGAGAGAA	DOCOTAGODA	CAGACGCTRCC
5601	OCCAGCION	GOOGTGAGTA	CICCCICTGA	AAAGCGGCA	TEACTTETES	GCTAAGATTB	TCAGTTTCCA	ANACCACCA	DOATTICATA	MCACCAGO"
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6001	GTAGGCGCTC		CAGAGGCGGC	COCCCTACACO	CCACCAGAAT	GATGATAGAG	CGTCTACCTO	CONCINCENCE	occocatcha	CCTCCACCACT
	CATCCCCCAB		GTCTCCGCCG	GCCARDANGERC	CCTCTTCTTA	CUCCUALCC	CCAGATCGAC	CCACACACAGO	CCCCCANAC	OCAGGTROCCA
6101	AAAGACCCCG	DOCADENDO	OCCUPICAN	GTASTCTATC	THECHTECH	GCAACITCTAG	CCCTCCTGC	CATOCOCOGO	COOCANDCCC	GCCC-TCG-TA1
1	7775700000		COCOCAGCTT	CATCAGATAG	ANCOTACCAA	CULTICADATE	CCARACGACG	GTACGCGCCC	accorneaca	CCCCACCATA
6201	COULTCAGE	8	TOCCATOCKY	TRACTITACE	CCONTRACTOR	CATCCCCCAA	ATCICUTANA	CCTACAGGGG	CTCTCTGAGT	ATTCCAARA"
	CCCAACTCAC		ACCOTACCC	ACCENCTUDE	GCCTCCGCAT	GIACGGCGTT		GCATCTCCCC	CACACACTCA	TARGETTETA
6301	ATOTAGGOTA	OCATCITICCA	CCCCCAINTIC	TRACCACIAC	GTANTOMAT	AGTICOTOCG		GAGGICGGGA	CCGAGOTTOC	TACCOCCO
	TACATCCCAT	CGTAGAAGGT	OCCUCATION	Accocacate	CATTAZ:ATA	TCAAGCACGC		CTCCAGCCCT	GGCTCCAMCG	AHALLIAL
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	COCAGINICOT	GCTTCCTCCG				-			ATTOORGANGE	
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10/0	CCTTOCCATT	CICCOAICOT	ACATCTTGAC	CACTOCCOS	ACCATCCGT	TUTTAGGGAA	AAGATCACCCA	TCGCGCATAC	CGACGCCCC	CAAGGCCTV1:
6801	GAGGTGTGG	TGACGCGAAA	COTOTOCCTG	ACCATGACTT	TGACKTETACTES	CATAVACTTC	TCAGTGTCGT	COCATOCOGO	CTCCTCCCAG	ACCATITITICA TCGTT:TTTCA
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1060	COCACOCOA	AMACCITICO	CCTAMACCGT	CCCCCTTCCA	CTGTAGCAAC	TK:TCATAGA	AARRONGCOOC	TCCCTATTTC	MCGCACACT	ACCCCTTCC!
7001	TCCCGGCACC		TOTTAATTAC	CHGGGGGGGG	AGCACGATCT	CCTCAAAAACC	CANCTACANC	TCCCCACAA	TOTABAGTITC ACATTICANG	CANGAAGCG" OTTCTTY: CC1:
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7101	CCCTACGGGA		GTTAAAAAT	TCANGGAGGA	TCCACTCCAG	ANGRECECTE	GACTCGGGCA	CCACACTITIC	CCCCCTCAGA	COTPICTACTIC:
7201	CENTROGANGE		CTCCACAGGT	CACHROCCAT	TAGEATTIGE	ACCITCUTCAC	GAMAGMEET	AMCTOGCGA	CCTATOCCCA	TITITICIO
	CCACCTTCO		GAGOTGTCCA	MOCCOGGIA	ATCGTAAAGG	TCCACCAGCG	CTTTC.CAGGA	TTTOACCOCT	GGATACCGGT	MAAAAGACC
7301	COTCATCCAG		OCCOUNT	TICCCAGCCO	TCCCATCCA	CCTTCCCACC	TAGGITCITCGC	OCCIOCAGICA	CTAGAGGCTC	ATCTCCCCC
•	CCACTACOTC	ATCT	COCCAGANC	MGGGTCGCC	AGOCITAGETT	CTANIXICCCG	ATCCARACCG	CACCGRCMGT	פשורורכשים	TAGAGGGG
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	CCCTTANACT		ACCIACLCAM	רנפעררער	COCCUSATION	CCSAGCTTGA	Transancate	GCCCAGATOO	GAGCTGTCCA	TOGICIOON
7801	GOACCACCAC	GCCGCGCGAG	CCCAMGICC	TCTACAGGG	CRETECCOCA	GCCTCGAACT	ACTUTTIGTAG	COCCITCTACC	CTCGACAGGT	ACCAGACCTC
				Pst						
7901	CHCCCGCGGC	OPCAGGTCAG		CTCCACATT	ACCTCCCATA	GACCICCAG	GOCOCOAGCT	AGATCCAGGT	CATACCTMAT	AAGOTCCCCO
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CCACTORGCC	CAAAAATTOC	TACCTOGACA	TCGGACGCT ACCCCTGCGA			CCGGCCGGAC		GCAGTCCTCC GGCCTGGCG CCCGGAACCGC	-	TCCTCCTGAA	GGAGATPAAC CCTCTAATTG GCGCTGGAGC CGCGACCTCG
TTAACGGTCT AATTGCCAGA	GGTATCCCAC CCATAGCGTG	: >	TGCTCCATTGG ACGAGGTACC	CTOTTOGATA GACCACCTAT AACCCAGGGG		•	AACGAAAAGG COTCCTACCG	-	• •	AACGACGCGC	CCCACTICAT CTTTTTANGE CTANGATTCG CGAACATTCG
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TAAA	TAAAAAAGGT		GTTCCGGACG	TUTOSCATIT	CCACTCGFTC	CGAMAGETETE	אניייייייייייייייייייייייייייייייייייי	רושוי איניניני		
	CONTRACTORICE		TOCTGACGC	CAACTORICA	CTGTTATTCC		מיהכדוראכם	CACAGTGGCA	CCAGICACON.	CCTOTOTOTATE
			ACGACTGCCC	CTTCAGCGCG	CACAACGACG	ACGATETATCG	CHRESTANGINE	כוניוניערנפו	CUCACAGO	
		Service .	CHEACTER CASE		ARGUSTATOT	CATALTGACATAT	NUTTICCARG	AGATTACAAD	TOTCAGCCGC	הנוצר הנאינה ממתמים הדרות
	CTACCACTOR				TCCICCTACA	CCTIXCTCGTA	TEMMOCIFIC	TCTAATGTTC	President Control of Control	רסכמשרורה
						CONTRACTAGA	AGATECECTE	OTTOCACACIT	TTAMCACC	ACCARTAGET
MOGN	MACAL	AGGAGGACAC GOOCAGCTG	CACACAMOLIC	ATTICATION			TETAKKAGA	CACGTGTCA	ANTHOROGO	אככיאכניאיניאינ
2	1	recreetara eccaredane					CARCITICANO	CTURACATOR	הכמכמכה	CATCEAACTE
		CATTITIOCOC TACOTACACAC	TCTTTCACTC			CULTATTRACIO	CHECKLACTORS	GACKTIVETACT	GACACACGTT	GFACCTTOGG:
,										

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12901	COCATOTATO		-	AACTUCITAA	TEXTACTE TACTE	מי אדרישינא נו	مدنعددشيي	ACCCCTARTA	TITCACCAAT	CCTATCTTCA
	CCOTACATAC	GGAGTTTGGC	COCCANATAG	TTKXX;CX;A1-T	AC'C'P':ATCAA	TENEDUS AT	נטטטטטטטער	TYRICACTCAT	MAGGGTTA	CENTAGAACT
13001	ACCCRICACTO	OCTACCOCCC	CCTCATTTRET	ACACCCATAY:	ATTRICARATE	CURCACATER	ACGATCKIATT	CCTCTCGCAC	GACATAGACG	ACACA: YSTICITE
	TOCOCCITOAC	: CGATGGCGGG	GOACCAAAGA	דידונות הכיניכים	TAAGCTCCAC	המתרותה	TKICTACCTAA	GGAGACCCTG	CTOTATCTGC	TOTOCCACAA
							- {	Hingiti	•	
13101	THECECOGOAA	CCGCAGACCC	TOCTAGAGET	נאנאעכאמנוני	じろいことのことと	ACCIONATION	CCCAAACGAA	GCGAAAGGAA AGCTTCCCCCA	GOCCAAGCAG	CPTGPCCGAT
	AAGGGGGGT	· occorcios	ACCUATCTCAA	CGITGICGCG	والإنطاء والعلاز	Truccentry	CUCTITICCT	TCGMOGCOT	ccocricore	GAACAGGCTV.
					I linititi					
13201	CTACACCACTO	COCCCCCCCC	OTCABATOCT	AGTAGCCCAT	TTCCAMICTT	GATAGGGGTCT	CTTACCARCA	CTCCCACCAC	2020222222	CTCCTCTCC:
	DATCCCCGAC	: מככמססמכסכ	CAGTCTACGA	TCATCOCCTA	AAGCTTCCAA	CTATCCCAGA	GAATGGTCGT	GACCOTOCTO	000000000	CACCACCC
			A.	_						
13301	ADCACCACTA	CCTANACAAC	_	תכטבות ביוני איבנים כאמנים	CGAMANAMC	ניומכביובנימ	CATTROCCAN	CAACGOGATA	GAGAGCCTAG	TOCACAAMAY.
	TECTECTEAT	· COATTHOTHO	AGCGACGACG	TOBOCGTOR	Germanns	מאכהאהאהיככ	GTAMAGRETT	GITGCCCTAT	CTCTCGGATC	ACCT: TTCTA
13401	CACTACATOG	* ANGACOTACO	COCAGGAGGA	CACCOCACGING	CCAMPICTICC	GUCCATCCAC	CCOTCGTCAA	AGGCACGACC	GTCAGCGGGG	PCTGGTGTGT
	CTCATCTACC	TICTOCATGC	acercencon	GICCCTCCCAC	ממדוירוראממ	STORY THE STATE	GOCAGCAGTT	recerocroa	CAGTECECEC	AGACCACACT:
13501	CACCACCATO	ACTCGGCAGA	CCACAGENCE	GPCCTGGATT	TREPROGRAMA	TACARCEES	TTRECACC	TTCGCCCCAG	GCTGGGGAGA	ATCTITTAAA
	CTCCTGCTAC	TOAGCCGTCT	GCTGTCGTCG	CAGGACCTAA	ACCETECKTC	ACCOUNTAGE	NAACCTCC-TC 20	AAGCGGGTC	COACCCCTCT	TACAMAITT
13601	ANANAANAA	GCATGATGCA	AAATAAAAA	CTCACCAACK	CCATTACACC	GAGCCTTRGGT	TITICITYSTAT	TCCCCTTANT	ATCCCCCCCC	COCCEATGTA
	**************************************	COTACTACGE	TITATITIE	GAGTGGTTCC	GGTACTOTOG	CIVICAACCA	ANAGAACATA	AGGGGAATCA	TACGCCGCGC	OCCOUTACAT
13701	TOAGGAAGGT	כבוכבובבב	CCFACCIACING	TOTOGREADE	CENTRICEAR	TRANSCORGE	CACHAGGITACT	CCCTTCGATO	CTCCCCTOGA	ccccccnrr
	ACTCCTTCCA		GEATICETET	ACACCACTCG	COCCOCOGIC	ACCOCCOCCO	CCACCCAAGA	GOGAAGETAE	GAGGGGGACCT	GOCCOCCAA
		Kyns								
13801	oraccrecoc		OCCTACCORG	CHICAGAMACA	CCATCCGTTA	CTCTGAGTTG	GCACCCLTAT	TCGACACCAC	CCGFOTGTAC	CTGGTOGACA
	CACCOCACOCO	CCATGGACGC	COGNICOCCC	CCCTCTTIGE	CCTAGGCAAT	GANACTENAC	CCTCCCCTATA	ACCINCTURETO	CCCACACATO	BACCACCTGT.
13901	ACANGTCAAC	COATGTOCA	TCCCTTAACT	ACCAGAACGA	CCACACCAAC	TTTCTGACCA	CCGTTATTATA	AMCANTGAC	TACAGCCCGG	GREAGECAAG
	TOTTICACITIC	CCTACACCGT	ACCCACTTCA	TRANCHISCT	GENERICETIO	AAAGACTTAST	CCCAGTAAGT	TITICITACTO	ATGTCGGGC	CCCTCCCTTA.
14001	CACACAGACC	ATCANTCTTO	ACGACCOGIC	GCACTCASCASC	CECCIACCIVIA	AAACCATCCT	CCATACCAAC	ATGCCAAATG	TCAACCAGTT	CANCILITACE
	GROTOTOTO	TAGTTAGAAC	TGCTCGCCAG	כנינופאכבבבם	CCCCCCCCCCC	TTTKKTAKTIA	CGTATCGTTG	TACOGITITAC	ACTIGCTICAA	GTACMATEM
14101	AATAAGTTTA	AGGCGCGCGT	GATOCTOTO	COCITIOCCTA	CTAAGGACAA	PUNCTITIENG	CTGAAATACG	ACTCCCTORA	OFFICACGCTG	CCCGARAGEA
	TTATTCAAAT	TECHOCOCCE	CTACCACAGC	OCCANCOGAT	GATTCCTGTT	AGTCCALY-TC	GACTITATGC	TCACCCACCT	CAAGTGCGAC	GCC-TCCCGT
				ţ	Paris			•		
14201	ACTACTCCGA	GACCATCACC	ATAGACCTTA		CATCHINAM	CACTACTTCA		-	_	ACCINCATOON
	TOATOMOOCT	CROSTACTOS	TATCTICAAT	ACTIVITIES	CTAGE:ACCTC	CHEATGANCT	-	-	CAAGACCTTT	CONCINCTAGEOR
14301	OCTANGELLE		ACTTCAGACT		وورسيدياله	CHITTHERE'NE	Ī.,		AAGCCTTCCA	TCCAGACATU
	CCATTICAAA	CTGTGGGGCGT	TOMOTOTON	נת:אאריםם	נאמדאהארצאני	ניאטאאטאט			Treathmast	M. M. T. T. T. A. I.
14401	APPENDENCE	CAGGATGCGG	CONTRACTOR		פניב בני אינא:אא	CENCIMONO			CCAGGAGGGC	TTACCATCA
	TAAACCACG	GTCCTACCCC	CCACCTGAAG	TOXIDENCE	COCACTOGIT	מאכאונכנים	TACKACCITACC	CCCTTCCCAA	BOTC TCCC	MATECTAGE

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CCCCACTOTOR CCCCACTOTOR TOCCATTOR ACCATACAT GAGAAACCT CTCTTCGRA CTCTTCGRA	GCAGI'TG'T'A CGTC(ACCAT	CTACTRESTC!! GATGACCA!C CACTCCAAGA OTGAGGTTC!	AAACCCC()	AGTECAGETA TEAGGTECK'T TTTTGAGEAA	AGANCHICHT ACCAACACCC TGGTFGTCA?		COTGCCCOC		ANGAGATICT TTCTCTACGA
GAACAGGGCG CTTGTCCCGC ACTTGCTAATA ACTTGCTAATT ACCCGAGGTC TGGGCTCCAG	ACCCAGTACC TOGGICATOO	CCCTCGTCCA GCCTCGTCCA GTTGCCCGTG CAACGGGCAC	GAGAACCAGA CTCTTGGTCT	GCATCGGAGG CGTAGCCTCC GAGCCGCACT	CTCGGCGTGA AAGCGCTCCG TTCGCGAGGC	GGTAGCTACG GGGCTATGCT CGCGATACGA	CCCCTCCAG	CCACAOCAGE GGCGTCGTCG CCGCCCCCO	GGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG
AGATGACAGG GTGAGAGAGA GACCTCCTGT CCCCTCCCGGAGGCGAGGGCGAGGGCGGGGGGGGGG	CAGCACCTIC GTCGTGGAAG	ACCTGCGGCT TOGACGCCGA GCGCCGAGCT CGCGGCTCGA	TCGCT FFCCC AGCGAAAGGG	CTGCGCAACA GACGCGTTGT GCGTCCTATC	COCAGGIATAG COCOCCAAG GCCCCGGTTC	GTCGATGACO COGGAGCCCO CCCCTCGGGC	CCTCCTTAAC GGACGAATTO	CCATCCCCCAC CCCTCCCCCAC	מאכאכטכטדט הדככאאמכטכ כאמהדוכסכמ
CCACGCTTGAA GCTCGAACTT AAATGCATYGGC GCTYGGCCCCGC GCTYGGCCCCCGC	TAARCAATGA ATTECTTACT	FCCTGACTTA ARSACTICAT CCGGTRCTYG GRECACCACC	ACCHOTICAA TOCACAAGTT	GACOCTACCO CTYCGATOGC GTCTCGCCGC	CAGAGCGGCG AGATGTTTGG TCTACAAAGC	GCGCACCACC CCCCACCACC ACCCTCATAC TCCCACCACC	SCCIOCUECE CONTRACTOR	GTCCAGGCGA CAGGTCCGCT CTGCGCGTGC	ACCCCCCACG ACCAACCTAT TCCTTCCATA
GRETACIONAL COSTATATA CONTITATO GRESALTATO MATTACITA	TACAACCTAA	PISCITITICAC ACGANACCITE CAPCANCTIT GIEGITIGANA	TETETGACEE AGAGAETGGG	CAGATCACGG GTCTAGTGCC CCTGGGCATA	CHACCCGTAT TTCCCAAGCA AAGGGTTTCIT	OCCUSTACTOS COCCOTONCO COCCATICAG CCAGTAAGTC	התכהיאסהת בתספדדסבה	TECCECCECAG ACTADORITIC GETTACICAT	
CHANTERAC CCTACACITY TICAACITY ACCTTYCCESS ACCTTYCCESS TCCCCCTTXS	CTTTCCGTCA	TCATGRACC AGTACTAGG CGCGCTCTA	CCAGTTTACC GGTCAAATGG	CCTACTCTCA GCACCAGAGT TTTACAAGGC	AAATISTTCCO RXXCTGCGC CCCOGACGCG	CACAAACGC GAGTTTGCGC CAGTTGACGC GTCACCTGCG	CHARTACE CICCOTOACO	ATKITCALTG TAACAGTKIAC TOCGLGACTC	ACCACCCACG ACCACCCACG
CCCCACHETT GCCGTRACAA GAGAGAAC CCTTCTCTTG AAGAGACGAC TTCGCCCGAC	MATACAGICAN TECTGTEGIT	COCHATICGG GCCTTAGACG THCCGCTACA	AACTCATCGG	TGAAAACGTT ACTTTTGCAA TGCCCCTACG	ACACAGGTTG TGTGTCCGAG	CTAXACCEC GACCTICGCAC CCAGAGATICA GUTCACAGGT	COCCCCCAACC	CCGGCGCGCA	
GCTANCATTC CCATTUTANG GCNT/GCCCCCC CGTCGCCCCCC CCGACTCCTC	CCCCTTACAG	ACCCTCAGAC TGGGAGTCTG CCCCGTGACC GGGGCACTGG	GTCTACTCCC	CCACCCTICAG GOTGCCAGTC	TGCGGCCTGG GCCAGCAATA GGGTCGTTAT	ACCOCOCOCC TGGCGCCCCG CACGCCCCCA GTCCTCCCGT	COTCACCACC	CTCCAAGGCT	CACACCCGITO GACTCGTACT CTGAGCATGA
TCTGGAGGGT AGACCTCCCA AACAGCACAC TTGTCGTCAC TTGCCACACAC	CCACTAGITT	AACTACOOCO TTUATOCCOC TOATOCAAGA ACTACOTTCT	OCTICTACIA COACCAGGCC COAAGATUTT GCTOOTCCGG	CCCACCATCA GOOTGOTAGT	GACTACACAC CCTTATATAC GGAATATAGC	COCOGOCACT GCGCCCGTGA ACTACACGCC	GCGCGCATCGT		CTCMGGGTCG GAGTCCCAGC ANACTACTTA TTTGATGAAT
CCTACGATGA GGATGCTACT AGGIGGCAGC TCCGCCGTCG GGCGACGCCT CCGCTGTGGA	ADMODANCE OCTONICANA TETTICITATES CENCENCITY	CCTTGCATAC AACTACOGCO CCALACOTATO TTCATCCCGC TTCCCAGACA TCATCCAAGA AACCOTCTCT ACTACCTTCT	OCTTCTACAA CGAAGATGTT		CACTGOTAAT OCATGTCCAT COTACAGOTA	ACTICCOCOTO TEACOCOCOCAC GAGGCOCOCOCAC	GACCOCTCAG	ACOGGGGGCC	AGGETATION CTCI TECCARGAM ANN ACCTICETT TETT
14501	14801	14901	15101	15201	15301	15501	15701	15801	15901

Figure 15J

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		Bigil	- T							
16101	CCADOTCATC O	GCGCCGGAGA	TCTATOOCC	CCCGAAGAAG GAACAACA	いいくしょくしょくしょう	ATTACAAGC	CTGAAAGCTA	ANGCGGGTCA	ANAGANAA	GRANGATICAT
	ODICCAGTAG COCOCCTCT AGATACCGGG GGGCTTCTTC	COCOGCCTCT	AGATACCGGG	OCCUTATION	CTTV TVICTO	TAATKITTECKI	CKICTTTCGAT	TINGCCCAGE	menum	CTTTCTACT.
								Sall		
16201	CATCATCANC 1	TTOACCACCA	COTTCAACTIC	CTYSCACGCTA	נג עשר הארכיב אני	הההארנאיהדא	CACTOGAMO	GITCHACCOCT	ANACOIOTT	TTGCSMCC:
	CTACTACTIO A	ACTOCTOCT	CCACCTTGAC	GACCITCCGAT	CANCINGENE	CCCTGCC17AT	GICACCTITIC	CAGCTGCCCA	TTTTGCACAA	AACCCTGG:)
16301	GCACCACCOT A	AGECTITACO	CCCCATTOAGC	GCTCCACCCG	CACI:TACAAG	CACGINITATO	ATCACCTOTA	CHOCCHCUAG	GACCTGCTTG	ACCAGCICAA
	_	TCAGANATGC	NGACCACTC9	CGACCTCCCC	CITURATETIC	GUTCACATAC	TACTCCACAT	OCCOCTOCTC	CTOGACGAAC	TCGTCCGGTT
16401	COARCOCCTC	GOSCINCTITIO	CCTACOGNA	REGGENTANG	מאכיאיזינים	רהדדינומבד	CCACCACTEC	ANCECANEAC	CTAGCCTAAA	OCCCGTAACA
		CCCCTCAAAC	COATCCCTTT	COCCGTATIC	CTGTACGACC	GCANCOBOTA	CCTGCTCCCG	Trecerrete	GATCOGATTT	COCOCATTICT
	154									Kpnt
16501	MON	TOCTGCCCOC	OCTROCACCO	TCCGAACIAAA	AGCGCCCCCT	AAAGETEEGAG	TCTOSTVIACE	TOOCACCCAC	COTOCAGCTO	ATTOTACCE
	DACGICGICC I	ACGACGGGC	CCAACCOTCCC	AGACITICITIT	Tenchecut	Triciacete	AGACCACTGA	ACCGTGGGTG	GCACOTCOAC	TACCATURGE
16601	AGCGCCAGCG A	ACTOGRAGAT	GICTIGAMA	ANATOACCGT	GGAACCTRAK	נידואטאבככם	AGGTCCCCGT	GCOCCCAATC	ANGCAGGTGG	COCCGGGACT
	TOCCOGICGE 1	TOACCTICTA	CAGAACCTTT	TTTACTGGGA	CCTACGACTC	CACCTCGGGC	TCCAGGGGA	COCCOUNTAN	TICOTCCACC	GCCCCCTGA
16701	OCCUPACAG 1	ACCOTOGACO	TTCAGATACC	CACTACCACT	AGCACCANTA	THRECACCRE	CACAGAGAGGC	ATCCACACAC	AMCGICCCC	GGTTGCCTCA
	_	TOCACCTOC	AMOTETATES	CIGATORICA	TCGTGGTCAT	AACGCTOCCG	Grencicco	TACCTCTOTO	THYOCAGOOD	CCAACGGAGT
16801	acadraacaa 1	ATOCCOCOOF	CCACCCCCTC	achistosacco	CGTCCAAGAC	CTCTACCAAG	GTISCAAACTRO	ACCCGTOGAT	GTTTCGCGTT	TCAGCCCCCC
	COCCACCOCC 1	TACOOCOCCA	CONCERCENC	כפעכפכבמים	CCACCTICTG	CAGATGCCTC	CACOTITINACE	TOGGCACCTA	CANAGCOCNA	ACTCCCCCC
16901	ממכמבכנימכמ	CCOFFCCACO	MAGTACTICC	CCCCCAGCGC	GCTACTOCCC	DANTATOCCC	TACATCCTTC	CATTGCGCCT	Acceceder	ATCGTGGCT
		OCCANGETEC	THEATGCCOC	GOCCOTCACO	CCANTISACCOC	CTTATACGGG	ATCITAGGAAG	GTAACGCGGA	TOCOCCCOA	TAGCACCGAT
17001	CACCTACCOC C	CCCAGAAGAC	DAGCAACTAC	CCGACGCCGA	ACCACCACTG	מאכניכפניכם	כבנאטטשבפב	CETCOCCAGE	CCGTGCTTGC	CCCGATTICC
		GOOTCHICTO	CTCGTTCATO	GOCTGCGCCT	TRIGHTECTICAC	CTTOTOCOGC	DOCORCYOCG	CCAGCGGTCG	GGCACGACCG	COCCIANAGO
17101	GTCCCCAGGG 1	TOCCTCCCCA	ACACACACACA	ACCCINGING	TOCCARCAIC	PROCTACCAC	CCCAGGATATCG	TTTAMAGCC	CONCIPIENTS	OTTENTIOCAG
		ACCEAGCCCT	recreated	TRACCACO	ACCASTIGICS	CACGATGGTG	GOCTCGTAGC	AAATTTTCGG	CCAGAAACAC	CAAGAACGT
										Sphi
17201	ATATOGCCT C	CACCTOCCOC	CICCGITICC	COCTCCCOCC	ATTICKGAGGA	AGANTGCACC	CTACATACACI		CACOOCCITOA	COCCUARCAIL
1		GTOGACGGCG	GACCCAAACC	שכבאכטפבכב	TAMORICITECT	TOTALCETOR	CATCCTCCCC	GTACCOGCCO	OTGCCCGGACT	GCCCCCCTA
	ā				Sphi					
17301	acorcoraca caccaccaac	CACCACCOOC	GOCOOCCCCC	GTCGCACCGT	CCN, ATT ACTOC	CCCCTATCCT	פנכניכונכונו	ATTCCACTOR	TCCCCGCGGC	CATTGACCALC
	COCAGCACOC 0	orcorocco	CCCCCCCCCCC	CAGCGTGGCA	OCCTACOCYC	CONCATARRA	CCCCCACCAA	TAAGGTCIACT	AGCGGCGCCG	CTAACCCCC
17401	GTOCCOCCA 1	TTOCATCOOF	OCCUPACAO	CCCCAGAGAC	ACTRATTAAA	AACAAGTERC	ATCTCACAAA	ATCAMATA	MAGICTOGA	CHCTCACGIT
		AACGTAGGCA	CCCCANCGTC	COCCIECTOR	TOACTAATT	THEFTEACG	TACACCTITIT	TAGITITATE	TTTCAGACCT	GAGACITCCCA
										ECHILA
17501	coerrooree 1	TOTANCTATE	TTGTAGAATG	THETHERATE GANGACATEA ACTITICETE TITTORICITE CGACACIOET	ACTITIOCGIC	הידטואיכנינס			CATCOCIAAAC	TRECAMENTA
))		ACATTGATAA	MCATCTTAC	CFICTOTAGE	TCAAACGCAG	AGACCTAROLIC		OCCCOOCA	GTACCCTTTG	ACCOPTICTA1

Figure 15K

PMRKAIISGOG MER682

GCAGGAAG?' CGTCSTTC(?) TAGCGGGAT! ATCGCCCCA!'	ACAGTGTCT* TOTCACAG**: TANAACCAN**: ATTTCGTTC**: CACTCAGCA*; GYTGGTCGT;	COCCCOTAG GCCTATGIN GTCCTATGIN CACCTACA '	AFFGFFCAAA ACTGCGFACT TGACGGCATGA GGGGCCCTA' CCCCGGGGAT	ACTITATITY: OGTATIATITY: CCATATITAT CTCAGTYSSTA GAGTCACAT	AATGATTACC" AATGATATA TTACCACTA! GTAACTCAC! CATTGATA!
ANDIACTATO GI TICTTOATAC CI CCTCTOGCAT TI GCAGACCGTA A	CCOGCACTC T CAGGAGGCAC T CAGGAGGCAC T CTCCTCCGTQ A CCCCCCCCTQ C GCCCCCCCTQ C		TCAGCCTGAN AGTCGGACTT AGTCGGACTT AGCCGTCCTA GGCACTCCTA GCCACTCCTA CACGACCTGT	GATCACCACA CCCTATACT CCGCAATAGA ATACCACATA	MCCCACACA TIGGORATIT ACCCCCACACC TCCACCACACC ATTAACACAC
TTCCACCGFF AAGGTGGCAA GATGGCCTGG CTACCGGAGC	AGCCTCCACC TCGGAGGTGG AGGGAGCATG GACCTGCCTC CTGGACGAGG		• • • • • •		r TCATATGCAN A GTATACGTT C CTACTGAGGC C CATGCCCACT C CATGCCCACT
AAAATTIKTS TITTAAAACC AAACSTGGTA TITCCACCAT	CCCTTAGAGG TATAGAGACC ATCTGCTCGG CGTAACGCTG GCATTGCGAC	CGCCGCGCGCGGGGGGGGGGGGGGGGGGGGGGGGGGGG	• • • • •		NT:TTACGAT TACANTGCCA TTTTTCTCAA AAAAGAGT ATATACTTA TATAAAGANT
AGCCSTAAT TYGCCSTAAT ATTTGAAGA TAAAGGTTGT	ACCEGECENT OFFICIALISA CACTAGGITT ACCATAGAGA TECHTINISS	COCCACTOTOR CONTECTION GITAGOSACT CIRCLOCTT GITAGOSCOM	TCC/NTTTCC ACCTTTCACG CCCTTTCACG CCCANCTCC ATGCCTTCCA	CCCCAAATCC CAAGTTAAGG GCACATATTT GCCCATATTTT CCGCCTATTTT	CHATGAAACC GTACTTTGG CHATACGTT CHTTACGTT CHAGACACTC
CTCGCTTTP37 GAGGGACACAY AAAGAGAGAAA TTTCTLYGTTT HIMIH	GTAVESTITION CAPTESTANCE AGANCTICES FOTTITIONS GTAS TEATICE CACCEACTORS	CACCARGEGG CACCATCRAGG TCTCACCGGTG ACCCCCCAC ACCCCCCACTGGGGGGGGGG	CCCCAGGCTCG CAGCCCGACC ACCCAGACT TGTACTGGAC ACACGACCTG	CCCAAGGTTG CGGGTTCCCCC ACCTACACTGA PCGTTCCACT ACCTAAATAT	AAGACTACC TTCTGATTGG AAAGTCAAGT TTTCAGTTGA FATAGAAAGG
TCARCTERODA AGTEGACICC GGATAAGTTO CCTATTCAAC	ANGATTAACA TTCTAATTGT CCGACMARA GACTGTCCCT GCCTACTGGA	GTTGTANCCC CAACATTGTG GCATCGTGGG CGTAGCACCC AGAGCTGCTG		CCCCTGRET CCGOOACCGA GAAGTAGACG AAGGTCAAAG TCCAGTTYG	
GRITGREGEET CCACCGCGGA MGATGETTANG TCTACGACTE	AGTGCANNT TCACGTTTTA CGTTCCCTCC GCAGGCCCG TCGCCCCAT AGCGCCCCAT	CTSCCCCTT CTSCCGCAAA ACACTGAACA TGTGACTTGT CGCCGCCAGA		CTCCCTACAA GACCGATGTT CAACGAAGAC GTTCCTTCTG ATAGGTGTCG	
CANTATGAGC GTTATACTCG ACACAGGCC TCOTOTCCCGG	CCAACCAGGC GGTTVGTCGG TOCCGAAAAG ACCGCTTTTC ACCGGTCCCA TOGGCAGGGT	TOCCAGOCCC ACGGTCCGGG CTCGCAAAGC GACCGTTCG GCGTCCATGT	TCGGCCGGCCCCCCCCGGCCCCCCCCCCCCCCCCCCCC		
FEMIN TCGACACCAG AGCCGTOGTC CTGGAACAGC	OTOGACCTOS CACCTGGACC CAGAGGGGCG GTCTCCCCGC CCTGCCCCCC CGAGGGGTCO	AMCCTOTOC TTTOCACACO CCAMTOGCAA GGTCACCGTT TGTCATOTAT	CATGCACATC GTACGTOTAG AGAACCCCA TCTTTGGGGT CGTACAAGGC	TTTAAGCCC AAAATTCGGG CTAGAAGAAG GATCTTCTTC TTACAAAGGA	MINITERECT COMMENTED COCCANGEN CCCOTTCON ACTTCACTC TGAACTCAGG
17601	17801 17901 18001	18101 18201 18301	18401	18701 18801 18901	19001 19101 19201

AND THE PERSON OF THE PERSON O		אדם סדי	TAN: Ann: CV : CCI :	נונדי נולא אדד	TAA ACC TOS CAA	7. YAN. 7. TTG	ACT
GOSTAATATO CCCATTATATO TCCATTGGITS AGGTAACGAC AACTTCCANA TTGAA: 1:1717 TGCTACAGAA ACGTACAGAA	MACATAGEC TTGTATEGY TGGCTCCCY ACCGAGGGGY	CCCCCCCTV CCCCCCCTV GCCCCCAGTV	ACTITION : CTITIONCO! GAMMITICE!	_	_	=	TCCMITANTT ACCTICATION
	CCTGTACTCC GGACATGAGG AAGCGAGTGG TTCGCTCACC	CCTTCTCTC CCTTCTCTCTO CCTTCTCTCTO	OCCAGEATTA COGTECTAAT ACCACTACT TGCTGTTCAG				CATCCCATTC
CTANTGRATT GATTACATGA CATACCAGCT GTATGGTCGA AMTCATGGA TTTAGGTACCT GAAAATGGAT CTTTTACCTA	GCACAAATTT CCTCTTTAAA CTACATGAAC GATGTACTTO	AACCACCACC TTAAAAACCT AATTTTTGAA		GOCCUTTGAC GOCCUTTGAC TOCCTCTATA ACCGAGATAT	_		CCCTTTOOCG
TTTATTOTT ANATACCA ACAGACCTT TSTCTCGAA GAATATTOA CTTAATACT AACAGSTCAG TTGTCCAGTC	-	CAACCCATT GTTGGGTAAA TTCTTTGCCA AAGAAACGGT		• • • • •		TCGCGTACTC TCGCCCACC ACGCGCGTTCG	TYLCKIATOOFA ACGCTAGCGT
TTMPASWAM ANYXXXTRTT AGACATATG CCACATGTA GCTCTACAAT TAMMACCTAA ATTTTGGATT	CANTETAAAT GTTAGATTTA GATAAGSCAA CTATTAGGSTT	TOTACAACGT ACCTGTTGCA GCCTCAGAAG CGCAGTCTTC		• • •	. ~ • •	AGAMACTICC TCTTTCAMCG TTGGCTMCCT	AANSTTRCFT TITCAANGAA
TACATHECTT ATGATTECA ATCTAMAGT CACCTATEA GTCCATACTA CTTACCANS GAATTATTCC	CCATCGAAAT GGTACCTTA AAAAATTTCT TTTTTAAAGA	CTTGACTATA GAACTGATAT ACATTCAAGT TGTAAGTCCA	* F S S E			CTCCTTCTTT CAGGAAGAAA TCTGGAATTTG AGACCTAAAC	ANTORECTE
CNGGCCTAAT GTCCGFAATTA AATTACHTTG TTACGACAACT TCCGACAACT TACAGAGACT ATGTCTCTA	MATMATTTE TTATTAMMC CTTCCMCGT GANGGTTCCA	ACGCTAGTCC TGCGACCAGG GTGCCCTTCC CACGGGAAGG	TTACATOST AATTOTACCA GGCCCACAC			ACCCCATGTA TYSCCTACAT ACACAACAAC TOTCTTGTTG	CAACTCACCA
CTATECECAA GATACGGGTT ATCCK: ACTTG ATCTGGAATC TACACCTTAG GTGTGATTAA	AAGAGTTGGA TTCTCAACCT AAGTACAGTC TTCATGTCAG	ACCTTGGAACTCG TGGAACCTCG TGGTCGCTAT ACCAGCGATA	ACCANGENTO PECTTECETAC PETTECECAT ACANGGOSTA	GCTCTACCCT CGAGATGGGA AAGAVAACCC TTCCTTTGGG	AGNAGATAGC TETTECACCO GGGTTACAAC CCCAATAGTTG	ACCTACAAGG TCGATGTTCC TCCTACACCA ACCATGTGGT	CAAGACCATA
GOCCAACANT CCGGTTGTTA CCGGCTTGTTA GCCCGGTTCT GCCCGGTTCT CATGAAAAAA CCACTGAAAAAA CCACTGAAAAAA	AAANTGAANT TTTTACTTTA CGACAAGCTA GCTOTTCGAT		GTGGAACTTC CACCTTGAAG TACGCCACCT ATGCGGTGGA		CACACCITTA OTGTOCANAT TTGACOGGGA AACTGCCCCT	TATCCCAGAG ATAGGGTCTC CAGGTGGGCA GTCCACCCGT	PECCETATE CULTIVIAGO ANGOCGATAC CEGNATATEC
AGANCTANTO TCTTGNTTAC GGTGTTCTGG CCACAAGACC ATAGAACCAG TATCTTGGTC TTACTGGTTT	TTTTCAGATA AAAAGTCTAT TGTATTTGCC ACATAAAGGG	OCTAGTOGAC CGATCACTG CGCTCAATGT GCGAGTTACA	ACACCTACCA TOTOGATOCT CATTTGCCTT	TATCTCTCCO ATAGAGAGCC CCTTCACGCG GGAAGTGCGC	TTACCTCARC AATOGNGTTO AAGCGCTCAO TTCGCGNGTC	AGGCFTC1A TCCCGAAGAT GGACTACCAA CCTGATGGTT	TECCCTATE AAGGGGATAG
19301 19401 19501	19701	19901	20201	20301	20501	20701	20901

Figure ISM

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					J. ij. iya adada	אירואיבניניני	ACGCCCT'AGA (CATCACTETY G		CCATCCACGA
21001			ACAGACT TGG	COCKET TOO				GTACTCANAL C	CTCCACCTAG 0	GGIACCTGCT
							ביינומאייות ו	ATCCAAACCG 1	TOTACCTOCO C	CACGCCCTT .
21101									ACT TOCACCIC G	GTGCGGGN 1
	COCOTOCCAA	GAANTACAAA	ACMACTICA	GMANC TEXT. A.C.	באואא, אבאו					IJAL
					1.0000000000000000000000000000000000000		GCTCCAGTGA	GCAGGAACTG A	AAAOCCATTG T	TCAAAGATCT
21201			ATMACAACC	AACACAACATT	THE THE PLAN.				TTTCOCTANC A	ACTITICTAGA
			TATITCITCS	The second	**************************************			accrocacca ?	TAGTCANTAC O	OCCUDENCY .
21301			TEXTECACION	Trincanticine	AAAGCTCCX.A	_		COGACOCOOT 1	ATCAGITATE C	CCCCCCACATA
	ACCAACACC	GOTATAMAA	ACCCCIOCAL	MC 1011COCC	CERTITION		CTCTTTOAGC	cermosent.	_	COACTCAAGC
21401	DAGACTGGGG	GCOTACACTO	CATCCCCTT	COCACCTION	CCCTCACTTT	-		-	-	OCTOAGTIC
	CTCTGACCCC	COCATONORER	CAGTCACTCC	TCCCCCTAG	COCCATTOCT	TCTTCCCCC0			AAGTCCACCC	AAAGCGTAC .
10017	Trrassition.		CTCAGTGAGG	ACCCCCCCATC	GCCCTNACCIA	NGANGGGGGC			_	. SUCESTICE DE .
11501	CONTINUE		GTGGACTATT	CHACTGCATG	THETECACO			ACTCCCATGG		GREGIACT!
10012	CCCCOOTIO		CACCTGATAA	GACGACGTAC	AAAGAGGTGC	GGAAACGGTT	GACCINGLAGIT			
		Kpre		•						
21701	CHARACTACO	•	CPCCATGCTC	AACAGTCCCC	AGGTACAGCC	CACCCTGCGT	COCMCCAGO	AACAGCTCTA	CACCITICATO C	CHEGEOGRA
10117	CAATAATGGC		GAGGTACGAG	TTOTCACCC	TCCATGTCCAG	GTCCCTACGCA	CLEVICON.			CTTTCAATAA
21001	Provide ACT		AGTOCCCAGA	TTAGGAGCGC	CACTICITIT	TOTCACTTGA	ANAACATGTA			GAAAGTTA1"
10012	GCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC		TCACGCGTCT	AATCCTCGCG	CTCACCACAAA	ACAGTGAACT	TITIGLACAL			COCATCOCTA
,	SCOT BARTIE		ACACTETEGG	GTGATTATT	אנככניבשכניכ	TRACCORCTO	COCCOTTTAA	AMICAMA		CCOTACCGAT
10617	TCCOTTTACO		TOTOACACC	CACTANTANA		AACCCCAGAC	CCCCANATT			TCACTCCACA
12001	TATACCACTO		OFTGCGATAC		-	NANCTCAGGC	MCAACCATCC		_	AGTGAGGTOF
	ACCCCCTCAC	COLCCC	CAACCCTATG	ACCACAMATC	~	THEAGTEE	2011201121			
									CGCGAGTTOC	CATACACACA
22101	GOCTGCGCAC	CATCACCAAC	-			AMGREGATION	PCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC			CTATISTICS.
		GTACTOGTTG	_		-		COCOUNTRACE	TOCOCOTOCA	GOTCCTCCGC (GTTGCTCAGO
22201	GFTGCAGCAC	TOGAACACTA	-			CCCTCTTCTC	CCTCTAGTCT	AGGCCCAGGT		CAACGAGTCC
	CAACOTCOTO	ACCTIGIDAT			_		THEFT	ACCOTAGIO	CATCAAAAGG	TGACCOTOCC
22301	OCCANCOCHO						AACGTGAGCG	TOCCATCACC	GTAGTTTTCC	ACTROCACOG
		A PATTOMANCC					CAGCETTEC	GCCTTCAGAG	AAGAACATGC	COLVACACITY
22401	COCTCTOOOC	GTTAGGATAC			-		CTCCCAAACG	COGNACA: TC	TTCTTGTACG	i-re-read
		CANTECTATO	TCCCCACCT	ATTTTCCCAAA	CTAGACTONA		III.			
			Shi				Section 2		ACTACATTIC GGCCCCACCG GTTCTTCACG	BFFCFFCACA
22501		GENERALARE TONTICOCCO	מאכאמינינפנ		פאבטאנים נאנינאניבידים	CONCRETE			CCGCCCTCCC	CNGANGTEC
1		COCCUPITIO ACTAACCOGC	CTOTOCOGCG		CAGCACOTASC GTCGTFGAAC	ניניאישרבאראא)		

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ATGCTTCCGT TACGNAGGCA	CAANSANTTE	CCAGGICTI-!	CISCIACAGOCITA GOGGOTOGO 1	OCACTTRICATE OCAGAACGCA	CCACTTICA XI	ANGAMANGA	GCAOTAGCCT	TGCAGCGC	AGTCAGCTCF	CCCCGCTTGA CCCCCCAACT	COTCCTGTTV3	CAGCOCCAGT	OTGOCOCOCA CTATCACATT GATACTCTA'S FCOMY	CCTCINTATO: GGACTATAR?
ATTINTCATA TAAATAGTAT	GICACCICTA			AGAAGGAGAA	GCCACCCAA	TCTTCCCOCG	CTCAGAAGGA	ACCARCCCC	CTAGTACCTC	CAGCTACCOTO	THEOTICS	GAAGCATCTG	CACCTATTCT GTGGATAAGA TGCTTGCCAC	CCCTCTCATA
CONSCIECT	ATCCTTCTAG TACGAACATC	CCCCCCACCA	GGTACTTGTC	TTCCCTCCCC AACCCACCCC	AACTAATCOT	COCCCANCC	CAGAACACTA	TOCAGGAGGT		CACCTTCCCC	TOTETECTAT		CTACGAACGC GATGCTTGCG GTGCCAGAGG CACGGTCTCC	TRECISEA DE ACRECECE
ATTICAATCA TAAAGTTAGT	TCCCCTCTC ACCCCARTAC	CAGCITACIAC	TTATCCACGT AATAGGTGCA	CACTITICCTC	AAACGCTACG AAACGCTACG	GGCGGGGGCT	CCHCCAGCCC	accer: tacto			CTCAGTACCA GAGTCATOOT		TCAGCCTTGG AGTCGGAACG CGTATTTACC GCATAAACGG	CAGCTGGCCT
CCACTGTACG	CGCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCTTCGCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCTTCGCCTTCGCCTTCGCCTTCGCCTTCGCCTTCGCCTTCGCCTTCGCCTTCGCCTTCGCCTTCGCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCTTCGCCCCCC	TOSTICANOST ACCACTÍCEA	CTTTAGATCG GAAATCTAGĆ	ACCOTANTITE TYCCATTANA	CONTRACTIC	CTCTCTCTCAT	CCACACGCGC	COCTCCCCCT	CCCCATTICC		ACGAGGACCO TOCTCCTRGC	CTACCTAGAT	ATAGCGGATT TATCGCCTAC ACTTCTAGCC TGAAGATGGG	ARCGGACAAG TCGCCTGTTC
CCTTTTKCALT	כאיזכיהאנאנו נזונאניאניוני	CAGAACAACG	TCAAGTTCGC ACTICAAGCG	CGRATTICATIC: GCCCAAGTAG	recartated Geotsacace	CCACCATTAC	CCCCCCCAC	CCTCCCCCCCC	CHICKCCACT	CGITTLCCACC	AGCGAAGACG TCGCFTCTGC	CCCTACCGCT	CCCCTCACC CCCCCCCCCCCCCCCCCCCCCCCCCCCCC	ACCGCAGCCA
מממנאלאיכ במכמיפאניא	CCCTCCCCAC	COTCACAAAG	CCCTCATCAA	GCACACTCAG	ATTCARCCOC TAAGITCAGCG	TCCTYXXTGT AGGARTGACA	ACCAGCTACC TCCAGCTACC	حددودعصود	COCTOCTCCT	TCGCCACCAC ACCGGTGGTG	AGGITTIGTA TCCAAAACAT	CCCCTCCTT	GCAGCGATGT COTCGCTACA CCAGCCCAAC	TRECRIFICEA
CTRICTTEAGE	TCGATTTK'AG AGCTAGAGTC	GCCCCATCAT CGCCGTAGTA	CACTTCCTCA GTGAACCAOT	GACACGATUG CTGTGCTAGC	CCAGGAGAAG	TTCTCTTTCT AAGAGAAAGA	PCCOCCCCC AGGCGGCGCC	CGANANACC	CCACCAMAGE	CCCTCTGAGT	AGCAGGACCC TCGTCCTGGG	AGTEGGGCOO TEAGECECOCE	TTOCAMONOC AACGTTCTCG ACGGCACATG TGCCGTGTAC	ACCCCTATCC TOGGGATAGG
TACTAGACTA	AAGCTCGCCT TTCGAGCGGA	C TOCAGGAATC	CCAGAGCTTC GGTCTCGAAG	CHCCCACGCA	COCCOCCACTO	GCGCCACATC	ANTOCCANA	COCCTCATCC	COCCOCTCOCO		OTCATTATCO CACTAATAGC	ACCAGGAACA	CTGCCACGCG GACGCTGCGC CGCCAAGAAA GCGGTTCTTT	TTTTCCAJA ACTOCAGAT AAAANGGTTT TGACGTTCTA
ATCTTCGCCT	MCAC	CAGGTACGCC	CATACOOCCO	CCATGCCCTT	CCCCATACCA	ACCATTTIGTA TGGTAAACAT	TCTTOGGCGC AGAACCCGCG	CTCCATACGC	OCACCOCOTC CGTGGCGCAG	AGAAGGACAG	CCTCCTCCTT	CCACAGGCAA	GCGCCATTAT CCCCGGTAATA ACCCCCCAAA TGGGGGGTTT	TTTTCCAAA AAAAAGGTTT
22601	22701	22801	22901	23001	23101	23201	23301	23401	23501	23601	23701	23801	23901	24101

Figure 150

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24201	CCTCGCTCAA	CCTCGCTCAA CGAAGTCCCA GGAGCGAGTT GCTTCACGGT	ATCTTTG	AGGGTCTTAG TCCCAGAAGC	ACGCGACGAC	AMBRESTER OF	CAAACCACTICT GTTTSCYCAGA	OCANCAGGAA COTTGTCCTT	AACAGCGAAA THGTCCCTTT	atgaaacitca Tact'ftcagt
24301	CTCTGGAGTG	TTGTTGGAC	TOTACCITCA	CACCICATA	CTAGCCTAC	TANAACTECAG	CATCGAGGTC	ACCCACTITO TIXXITONAAC	CCTACCCADC GCATGGGCCG	A cti aaccta Traatitgrat
24401	CCCCCCAAOO	PCATOAGCAC AGTACTOGGG	AGTCATGAGT TCAGTACTCA	CTCCACTANC			CTCTCCCTAC		AGNACANACA	GAGGAGGGC**
24501	TACCCCCAGT	TOCCOACGAG ACCCCTOCTC	CAGCTAGGGC	GCTCACTTCA CCACCCAAGT	AACGCGCGCTC	CCTCCCCCTCT CCACCCCCTCA	TREATHCACCE ACCTCCTCCC	ACCCAAACTA	ATCATCCCCC	CAGN:CTCT.T GTCACGAGCA
24601	TACCGTOGAG CTTGA ATGGCACCTC GAACT	TACCOTODAG CTTOAGTOCA ATOGCCTC GAACTCACGT	Greca Techeconst	CTITIGCTAAC	CCAGAGATAC	ACCGTANGET TCGCGTTCGA	AGACCANACA Tepectitus	TTCCACTACA	CCTTTCGACA	GOCCTACGTA
24701	COCCAGOCCE		CAACOTOGAG OTTOCACCTC	CTCTGCAAGC	TRGSTCTCCTA ACCAGAGGAT	CCTRGGAATT	TTGCACGAAA	ACCOCCTTOG TOCCOGAACC	OCAMACOTO COTITIGGAC	CTICATTCCA
24801	CGCTCAAGGG	COCTCAAGGG CGAGGCGCG GCGCTACG GCGAGTTCCC GCTCCACGCGCG GCGCTCATGC	COCGACTACO	TCCGCGACTG AGGCGCTGAC	CGTTTACTTA	TTICTATGET AAAGATACGA	ACACCTGGCA TETGGACCGT	CTCCCCCTAC	OGCOTITION CCOCAACCO	AGCAGTISCTT TCGTCACGAA
24901	GGAGGAGTGC	AACCTCAAGG	AGCTGCAGAA TCGACGTCTT	ACTGCTAAAG TGACCATTTC	CANANCTIGA	AGGACCTAFG TCCTGGATAC	CHOCCHO	AACGAGCGCT	CCGTOGCCOC	OCACCTOSICY: CGTORACCO!:
25001	GACATCATT CTOTAGTAAA AGCOCTCAGG TCGCAAGTC		CCTGCTTAAA GGACCAANTTT GCCACCTGCT CGGTGCACGA	ACCCHACAAC TRAGACGTTG GTGCACTTCC CACGTGAAGG	ACACCIOCA TECENOACCO TACCIACTIT ATECETGAAA	AGACTTCACC TCTGAAGTGG GTGCCCATTA CACGGGTAAT	ACTCAAAGCA TCAGTTTCGT ACTACCGCGA TCATGGCGCT	TOTTGCAGAA ACAACGTCTT ATGCCCTCCG TACGGGAGGC	CCGCTTTOOG CCGCTTTOOG CCGCCTTTOOG	AAATAAGATU GCCACTTACTA COGTGACGAT
25201	CCTTCTGCAG GGAAGACGTC		ACCTTOCCTA	CCACTCTGAC GOTGAGACTO	ataatggaag tattaccttc	19 to \$2	TCACCOSTCT'A ACTCCCAGAT	CTOGAGTOTC GACCTCACAG Psil	ACTOTCOCTO TGACAGCGAC	CANCCTATKIC GTTGTATALT
25301	ACCECOCACE	CENECCTOST CENECOCICA	TTOCARTICO			AATTATCGGT	ANTTATCERT ACCTITICACE THATACCCA TODAMCTCO	TOCAGGGTCC ACOFCCCAGG	CTCGCCTGAC GAGCGGACTG CACGAGATTA	GANNAGTECTI CTTTTCAGG" GOTTETAGGA
25401	COOCHCCOO	_	ACTCCCG000C TGAGGCCCCC	•		COCCETTANAC	AFFICIAL	GATOCTOCOS	GTGCTCTAAT TCAACAAAGC	CCAGGATATT
25501		MANCEANTCE COCCOCCTA TETECTAGO GEOGOCOGNT TITETECTAC GAAAGOTACO AAAGACGATO CTTTCCTGC	ATCCCTCGA TACCCTCGA GGGGGTTTTAC CCCCCAAATG	TACCACCACA ATGGCGGACT TTYGACCCCC	CACTAATYAGO ACTECCOGYCA TCACCCCCT	PCCCGATTGTA GGACCTCAAC CCTCGAGTTG	AGAACCGGTT CCAATCCCCC GGTTAGAAAA	AACGITCGGT CGCCGCCGCA ACGGCTACGT	AGTIGITICO GCCCTATCAG CCXGATAGIC	CANCARKES. CANCARKES. GRUGIKAGOS

Figure 15P

PHRKACTSgag Nerrenz

Pstl

				STATEMENT	*					
25701	GGGCCCTTGC	TRECEAGGAT	CCCACCCAA	AAGAAGCTCC	NATITATOR	נינגענגנענפ	עעטטעטטעט	ANTACTOCICA	CAMPCAGGCA	CARRACETTE
•	CCCOGGNACG	CCCOGGNACG ANGGGTCCTA	CCORNER	TTCTTCGACG	איזיטטטאנאע:עראר	CLYSTRAGING	CINCIPCOTOC	TTATGACCCT	OTCACTCCGT	CHICKLON .
						ferefill				
25801	TOCACGAGGA	GCAGGAGGAC	ATCATCANG	ACTROCANGAR	נינידאהאהיהאה	האאהכידיבככי	AGGTEGNAGA	CGTGTCAGAC	GALACACCUT	CACCETTORY .
	ACCTGCTCCT	CCTCCTCCTO	TACTACCITIC	TGACCCTICTIC	GGATICTGCTK:	CITTCGAAGOC	TCCAGCTTCT	CCACAGACTO	CTTTGTGGCA	GTGGGAGCC 'A
25901	CGCATTCCCC	Traccoacac	CCCASAAATC	GGCAACCGGT	TECACICATION	CTACAACCTC	CACTCCTCAG	0000000000	CACTGCCCGT	TCIACCIACCC
	GCCTAAGCCC	AGCGGCCGCG	CONTERTING	CCGTTGGCCA	ACCITCGTACC	מאזרויזואט	CCGAGGAGTC	crecococce	GTCACCOCCA	AOCOOCTOCO
26001	ACCOTAGAT	GOGACACCAC	TOGANCCAGG	OCCUSTANST	CCMACAGCC	CCCGCCCTTA	CCCCAAGAGC	AACAACAGCG	CCAAGGCTAC	CIXCTCATOG"
	TTOOCATCTA	cccrotosta	ACCITICACTOC	CGCCCATTCA	GSTICGTUGG	CRACARCANT	COCCITICACO	TIGHTOTOGC	GGTTCCGATG	OCCUMENTACO:
26101	OCCOOCACAA	GAACGCCATA	othecticet	TOCANGACTO	TYCOCCOCAAC	ATCTCCTTCG	CCCCCCCATT	TETTETETAC	CATCACORCO	TOXIC THEFT
	COCCOSTOTT	CTTGCGGTAT	CAACGAACGA	ACGITICTIGAC	ACCCCCGTTG	TAGAGGAAGC	GCCCCCCCAA	ACANGAGATO	GTAGTOCCOC	ACCTIFINACION.
26201	CCOTAACATC	CTCCATTACT	ACCONCATCT	CTACAGCCCA	TACTRICACES	CCCCCCACAGCCC	CAGCNACAGC	ACCOCCACA	CAGAAGCAAA	CHACKACCOCA
	OCCATIGIAG	GACCTANTGA	TCCCAGTAGA	GATOTOGGGT	ATGACGTYCC	CACCGTCGCC	Grearieres	TCCCCOOTCT	Greatena	בכנוכיוספרכזי
26301	TAGCAAGACT	CTGACAAAGC	CCAAGAAATC	CACAGCGCG	GCARCAGCAG	CACTACACTACC	GCTGCGTCTG	GCCCCCAACG	AACCCGTATC	CACCCCCGAG
	ATCOTTCTOA	CACTOTITICO	GGTTCTTTAG	GTGTCGCCGC	COTEGICATE	CTCCTCCTCG	CGACCCAGAC	cococomoc	TTOOOCATAG	CTOCOCOCTC
26401	CTTAGAACA	GUATTITICC	CACTETRITAT	CCTATATIFIC	AACAGAGCAG	GROCCMOAA	CANGAGCTGA	AAATAAAAA	CAGGICTICTO	COATCCCTCA
	GAATCTTTGT		GTGAGACATA	CCATATAAAC	TRICKER	CCCCGGTTCTT	GETCTCGACE	TITATITI	GTCCAGAGAC	OCTAGGGAGT
26501	CCCOCAGCTO		NAMGCCAAG	ATCAGCTTCG	GCGCACACTG	GNAMAGCGG	AGGCTCTCTT	CACTAAATAC	TOCOCICCIOA	CTCTTAAGGA
	GOOCGTCGAC		###COCIFIC	TAGTCCAAGC	כמכשוצכמענ	CTTCTOCGCC	TCCGAGAGAA	OTCATITIATO	ACCCCCCACT	CAUAATTCC .
26601	CTAGTITCOC	g	AAATTTAACIC	GCGAAAACTA	CGTCATCTCC	AGCOCCCACA	CCCCCCCCCA	OCACCTOTTO	TCAGCGCCAT	TATITACCAM
	GATCAAAGCG	8	TITAAATICG	CGCTTTTGAT	CCACTAGAGG	TCCCCCCTCT	OGCCCCCCOT	CCTCGACAAC	ACTCCCCCCTA	ATACTCGTTC
26701	GARATTECEA	5	GTGGAGTTAC	CARCCACAM	TGGGACTIGG	COCTGGAGCT	OCCCANGACT	ACTCAACCCG	AATAAACTAC	ATGARCCCC
	CTTTAGGGT	B	CACCTCAATG	greconstra	ACCCTGAACO	CCGACCTCGA	COCCUTICATOR	TCACTTGGGC	TTATTTCATO	TACTCGCGCC
					Ū S	ILIOCI				
26901	GACCCCACAT		GTCAACGGAA	TACOCOCCCA	CCCMANCCCA	ATTUTCHUNG	NACAGGCGGC	TATTACCACC	ACACCTCOTA	ATAACCITAA
	CTOCOCOTOTA		CAGTTCCCTT	ATGCCCCGGT	CAXCTITICOCT	TANGAGGACC	TTCTCCCCC	ATANTOGIGG	TOTOGAGCAT	TATTCCAAT
26901	TECEGORAGE	100000010	cccruststa	CCACCANACT	CCCCCCTCCCA	CCACTGTGGT	ACTICCCAGA	GACGCCCAGG	CCONNCITICA	CATCIACTANC
	ACCOCCATCA	ACCORCCAC	GOGACCACAT	GGTCCTTTCA	THE CHANGE T	OCTOACACCA	TCAACCOUTCT	CTGCGGGTCC	GOCTTCAAGT	CFACTGATTO
27001	TCAGGGGGG	AGETTOCOOD	COCCUTICGE	CACAGOOTTIC	מכתבטבבבנטט	CCACCCTATA	ACTCACCTGA	CANTCAGAGG	GCGAGGTATT	CARCTCAACT
	Agrececoed	TCOAACOCCC	GCCGMAGCA	OTGRECCACG	CCARCAGICC	CGTCCCATAT	TCACTCCACT	GITAGICICC	CGCTCCATAA	Greangray.
27101	ACGAGICOGE	GAGCTCCTCO	CFFCCFCFCC	GTCCCGGACGG	GACATTICAG	ATCGGCGGC	CCONCCACTC	TTCATTCACG	CCTCGTCAOG	CANTECTANG
	TOCTCAGCCA	CTCGAGGAGC	DAACCAGAGG	CAGGCCTGCC	CTCTAAAGTC	TAGCCACCAC	GGCCGGCGAG	AAGTAAGTGC	GCAGCAGTCC	GITACCATTC
	F.									
27201	TCTGCAGACC TCD	restections	Achedecte	TOGAGOCATT	CONVINCTOR			CCATCOGICT		CTICTCOCK
l I	AGACGACTOG AGO	AGCAGGAGAC		ACCTCCOTAA	CCTTGAGACG	TTANATMCT	CCTCANACAC	COTACCCAGA	TCAAATTCCC	GANGAGCCT

TTCACCTCTC COTCTCCTT*1 CCCGAGGATC ATATCTAGTG OGOCTCCTAG TATAGCTC** TAGTTGAGG GGAGAGGG*A ATCACTCGC CCTGTCCC** TAGTAAATAC AGAATTA**A		TTOTOGTOGO AGGNATOGNO TCANTACTC TOTTTACCAO AGTTATTONO ACNATOGTC	_	ANOGENCE COCENCENT TECCOLOGICA COCENCIA COCENCIA COCENCIA CACOCECCA AGROCIANCE CAGECOCOTTE	TATADARIC ACCACAGN ATATTTACG TGGTGTT'S GAGTATARIC TTACAGTT'S CTCATATTAC AATOTCAANA	AGTATAGOT GTOSCCCCCA TCATATTCAA CACCGGGGGGG CTATATTAAA TACAAAAGCA GATATAATTT ATGTTTTCC CTCCTTCCAA AACAAATTT'A GACGAACGTT TTGTTTTAV'T TCTATGTGGG ATATTCTC'A AGATACACCC TATACCTC'A AGATACACCC TATACCAGGT
ACTOANTOTT A THACTTACAN T CTTTGANTTO C GAAACTTANC C GOCCCCTOC OCCOCORO A				TTETETGGGT AAGAGAGGA CTAGGTTTAC GATCCAAATG	CCTCCTCTCT CCTCCTCACACACA TCACACTACA ACTCTCATCATCT	ATGAGGAAG TACTCGTTTG GTACCCTACT CATGAGATGA GCTTTACTCG ACAATGAGC AACAATTGAC TTGTTAACTG
GACCAGCTACG CTGCCTATGC ACTTTACTA TCAAAAGGAT GTTTACCCAG CAAATGGGTC				ATACTAACOC TATGATTCCO GTACATAATC CATGTATTAG	CCTANTRAGT CGATTACTCA CGATTACTCA CGGATCCAGG	TACCATGFAC ATGGTACATG CCANACCAGA ACCACTANCT TGGTATTGGT CANAGGGGAC GTAAGGGGAC
			CTACTGTCATO	CTTTATACTT GANTAAGAA AGATGATTAG TCTACTAATC	COCACCTGAN CCATCGACTT TATCCTATTT	TOTICGACAT ACACICCTGTA TACAGTECTE ATOTICACAA ACCATTACAG TECATTACAG TECATTACAG ACCAGTTATG
	GVANTGTAGT CVANTGTAGT TCTTCACCCG AGANGTGGGC	ACTOTOTOTO TGCTCTCTTG CTACCGCCTG	MANACCCAG TTTCCCCTC	THGTGATTCT AACACTANGA TCGCCACCCA AGCGGTGGGT	A ATGITACATT TACANTGIAA GIATCETGIT COTACGACAA	TITTATIGAMA MANATACTIT CTATICETAAT GATACGATTA TANCTIACAA ATTICAATITIC COTICATITIC COTICATITIC
CCTANCTTH GGATTGAAAG GCCACAAGTG CGGTGTTGAC GGGAGAAGTT CCCTCTCGAA	CCTAACCCTG GGATTGGGAC AAGGCCACCG TTGCGGTGGC	CACACTCAGA TIXCACCACAC ACGTGGTGTG	GTATTAGGCC CATAATCCGG	ATTCTCTGTC TAAGAGACAG AACGCTGGGG	CCAGCCTGTA GGTCGGACAT AAATTGGGAA TTTAACCGTT	ESTITUTE CIT TACTITICCA CAT ATCAVAGGT THE TECTOCACTG AND ACGACGTGA AND ACGACGTGA AND ACGACGTGA AND ACGACGTGA INT TANACCCCCC INT TANACCCCCC TANA TANACCCCCCC TANA TANACCCCCCC TANA TANACCCCCCC TANA TANACCCCCCC TANA TANACCCCCCC TANA TANACCCCCCCC TANA TANACCCCCCCC TANA TANACCCCCCCC TANA TANACCCCCCCC TANA TANACCCCCCCC TANA TANACCCCCCCCC TANA TANACCCCCCCCC TANA TANACCCCCCCCC TANA TANACCCCCCCCCC
	TTGCAACTGT AACGTTGACA CCATCCTGTA GGTAGGACAT	AACCCAGACO TTOGGACTGC CACCGGCCGC GTGGCCGGGG	AACCCTTAGG TTGGGAATCC	CCACCCCAA CCACCTTTTTA GTCGAAAAAT	TTTTANG AAATTC CACAAAA GTGTTTT	CTTTTAT GAMMEN TGECACT ACCOTON NGMMN TTCTTTT GAMTAGC
ACTATCCOSA TGATAGGCCT ACACCTGGTC TGTCGACCAG GGCGTCCGGC CCGCAGGCCG	TCACTGTGAT AGTGACACTA GCTCCTATCG CGAGGATAGC	CAACAGTTTC GTTGTCAAAG ACGAGTGCGT TGCTCACGCA	GAGCTTAGAA CTCGAATCTT Xbsi	•	AAAAGTUUA TTTTCCACCT GCTTATTCOC	ACCENTATE TEAGRAMCMC ACCENTIFICE TATTGMOGIA ATACCECT ATTACATA
CCTCCCGGCC GGAGGGCCGG TGCGCCTGAA ACGCGGAACTT CCCCGGGGCAC	CCCTGTGTTC GGGACACAG ATATACTGGG	CTOTGATTTA GACACTAAAT CCGGGAACOT GGCCCTTGCA	AACAGGAGGT TTGTCCTCCA	TCAGOTTACT AGTCCAAAGA TGCACATTIG ACGTGTAAAC	KENT GOTACCACC CCATGGTGGG ATGAAAAGCT TACTTTTCGA	CCAGOGTANA GOTCCCATIT CANANTIGE GACGENGET CTCCGTCGAA AANGTINGE FIFTEANTEG
27301 27401 27501	27601	27801	28001	28101	28301	28501 28601 28701 28801

PMRKAdSgag MENGBZ

. 28901	COCCTACAAC	CTTGAAGTCA	CCT.MUGALC	ATGTCAGGAT	CHEACTHAGE	CCAGCACCTG	TCCCCCCCCCACAT AGGGCCCCCTA	MCANGETCA ACANGETCA	OCTTOATOTO	CCACCCACIC CCT KOOTH HTS
29001	TAACAGAGAT	GACCA			CCATACTTACA	TUTACCACAA	ATACACCCCA	AGTITICTUCC		ACTOGGATAA
	ATTOTOTOTA	CHGGTTGTGT	TOOTTICCCC	CACCACCAATT	CCCTGAATG	אהאוזאיזאיזאיזא	TATCHICOCT	TCAMCACCC		
29101	CTTGGGCATG		_		Traicttaita	TTATCHART	CATCTCTCTC	CTABARCCCA	AACOCGCCCG	ACCACCCATC
	GAACCCOTAC	ACCACCAAGA	CCTATCCCCA	ATACAAACAT	ACCOUNTANT	MATACACTOA	GTAGACGACG	GATTICCCUT		
29201	TATAGTCCCA	TCATTOTGCT	ACACCCAAAC	AATTATTAAA	TCCATAGATT	שיאנגאיארידה	ANACACATGE	ACTIFICACT		-
	ATATCAGGGT	AGTNACACGA	TOTOGGITTO	TTACTACT	AGGTATCTAA	CCTCCCTCIAC	TITCTCTACA	ACALAAGAGA	ATCICATACT	AATITACICT
	•	Yhol								
29301	CATCATACCT	CATCATICCT CCACTITITIA			CTTTTTTTTC CONSCICENC	CONSCIECTAC	ATTOOCTOCG	Grincheach		
	. GTACTAAGGA	GTRCTAAGGA GCTCAAAAAT	ATAATCACTO	CCAACAACCC	האסאאראים הבאכהאההוא	הכאכמאממינה	TWCCGACGC	CAMAGAGTGT	AGCITCATCE	GACCIPAGG
					181		1			
29401	CCCTTCACAG	TCTAT		-	CACTCATCTA CACCATCATC	CACCCACATC	ACTIONCINCA	ACTIGICA TOCCITIAT	CCAINTGCATT	CITCACCITACA
	COCAROTOTC	MONTANACGA	ATTACTION	CAGTGGGR	פרויאין אין ארי	מורואוווויו א	Ecoff	-		
20501	and the second	Trickly after	AGACACTATIC	CCCAGTACAG	GGACAGGACT	ATAGCTGAGC	THUTTAGANT	PICTURGAT TOTTTABLER	TOAMTTTAC	TOTOACTT!
	CACACGCGAA				CCTGTCCTGA	TATEGACTEG	ANGMATCTTA	AGALATTAAT	ACTITAAATO	ACACTOMAN
29601	CHEST TO A PIPE				CCTCCAAGCC	TCAAAGACAT	ATATCATOCA	CATTCACTCG	TATATGGAAT	ATTCCAAGI T
	DACGACTAAT		TAGACCCANA		GGAGGTTCUG	AGITICIGIA	TATAGTACGT	CTAAGTGAGC	ATATACCTTA	TAAGGTTCAA
							Pel			
20701	CAPACAATEA	AAAAAGCGAT	CTTTCCGAAG	CCTCGTTATA	TCCAATCATC	TCTGTTATES	TGTTCTGCAG	TACCATCTTA	OCCUTAGETA	
40167	COATGITACT				ACCTTAGTAG	AGACAATACC	ACAAGACGTC	ATOGTAGAAT	COCCATCCAT	ATATAGGAT
1000	CALIFORNIA PER		-		CCAACTITICC	כבפכשכבנופכ	TATGCTTCCA	CTCCAACAAG	TTGTTGCCCGG	COCCUPING
***************************************	COAACTIOTAA			GTACTTGGTG	CONTICAMO	DOCCCGGGGG	ATACGAAGGT	GACGITOTIC	AACAACGGCC	AACAACGGC GCCGAAAFA 1
									X a);H
									Boll	
29901	CCAGCCAATC AGCCT	Agenteace	Accrictece	ACCCCCACTO	ANATCACCTA	CTITAATCTA	ACAGGAAGAGA		CCCTAGATCT	
	COTCOOTING	TCGGAGCGG		TROGRETICAL	TTTAGTCGAT	CANATTAGAT	TOTOCTOCTO	TACTGACTGT	GOGATCTAGA	TCTTTACCTO
1000	CHARTERIA		CETTOCTAGAA	AGACGCAGG	CAGCGGCCCA	CCAACAGCGC	ATGAATCAAG	AGCTCCANGA	CATOOTTAAC	THOCACCAGE
	CCTTAATAAT	_	_	1CTOCOTCCC	GTCGCCGGCT	correspond	TACTTAGTTC	TCGAGGITTCT	GTACCAATTG	AACTTICOTICA
10101	GENANAMORIA		CTCGTAAAGC	ACCCANGE	CACCTACGAC	AGTAATACCA	CCGGACACCG	CCTTAGCTAC	AAGTTGCCAA	CCANGCGT".
	COTTITICCCC		_	TCCCCTTTCA	CTOCATGCTG	TCATTARGE	GCCTGTGGC	GONTCCATO	TTCACCOUT	GCT PLOT ALL
10201	GAAATTOORO	GICATOOTOG	CAGNAAAGCC	CATTACCATA	ACTCAGCACT	CTRITACAMIC	CGAAGGCTGC	ATTICACTICAC	CITIOTICAAGO	ACCTGARGAT
	CTTTANCCAC			GTAATGGTAT	TCACTCCTCA	OCCATUTION	GCTTCCGACG	TAACTGAGTG	DACAGTICC	TOCACTOCTA
				Brita	ł					
30301	CTCTCCACCC TTATT	TTATTANGAC	CCTOTGCGGT	£.	TTATTCCCTT		MANAMATAA	THANGCATCA	CTTACTTAMA	ATCAGTTMIC
1	CACACOTGGG	GAGACOTGOG AATAATICTG GGACACCCCA	GGACACCCCA	GAGTITICTAS AATAAGGAAA	NATANGGATAN	ATTENTATT	TTTTTTT	ATTICGINGT	CANTCANTIT	TRITCAMICG

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30401	AAATTTCTGT	CCAGTITATE	CAGCAGCACC	rectrosect	CCTCCA(XCT	נחמנדאדופנ	AGGTTCCTCC	TOCCTOCALA	CFFFCFCCAC	ANTCTANATO
	TTTANAGACA	GCTCAAATAA	Greenestes	ACCAACCCCA	היאיימידנימאיאי	האתנאדאתה	Tremagned	ACCGACOTAL	CARAGAGGIC	TTALIATI I AL
30501	GAATGTCAGT	TACCACTOF	TCCTGTCCAT	CCGCACACCCAC	TATCTTCATG	THEFT AND A STATE OF THE STATE O	TGAAGCGCGC	AARACCETCT	CARCATACCT	TCAACCCCC:T
	CITACACTICA		ACCACAGGTA	CARICHANGE	ATAGAAGTAC	ANCINACIPIET	NCTITOTICOGG	TTCTCGCAGA	CTTCTATOOA	ACT TOCKSON A
30601	GTATCCATAT	GACACOGAAA	ccochectee	AACTGTGCCT	THETTACTE	Crecenter	NTCCCCAAT	OCCULTICARD	AGAOTECCCC	TOOGGT.ACT :
	CATAGGTATA	Characettt	CCCAGGAGG	THEACACOGA	ANGANTGAG	GACCCAAACA	TAGGGGGTTA	CCCAAAGITC	TCTCA00000	ACCCCATGAG
				Ξ.	Falts					
10701	PEPPICEUC	TATCCGAACC	TCTAGTTACC	TCCANTISCA	TRICTTOCOCT	CAAAATCAGC	AACREATER	CTCTGGACGA	GCCCCCCMC	CITACCTCC:
	AGNANCOCOG		AGATCAATCG	ACCTTACCGT	ACCIANCICCGA	GTITTACCCG	TYCCCCCCAGAGA	GAGACCTGCT	CCCOCCOTTO	GANTCCAG 1
LOBOL	AAAATOTAAC	CACTICTICAGE	CCACCTCTCA	AAMAMCEAA	GTTANATATA	AACCTOCAAA	TATCHOLACC	CCTCACAGIT	ACCTCAGAAG	CCCTAACTKIT
	TITTACATTO		COTOCACACT	THYTHE	CAGTTTGTAT	TTCKOACCTTT	ATAGACGTCS	CCAGTGTCAA	TOGRATICITIC	GOONTIGACA
10901		Ž	TOCTOCOCO	CAACACACTC	ACCATICAAT	CACAROCCCC	CCTAACCGTG	CACGACTCCA	AACTTAGGAT	TOCCACCCAA
1000	CCGACGGCGG	8	ACCAGOGOCC	GTTGTGTGAG	TEXTACCITA	GIGTCCGGAG	CGATTCGCAC	GTGCTGAGGT	TIGAATCOTA	Accordage !
11001	GCACCCTCA	S	ACCAMACTA	OCCUTOCAAA	CATCAGGCCC	CCTCACCACC	ACCGATAGCA	GTACCCTTAC	TATCACTOCC	1CACCCCC17
	CCTOOCOAGE		TCCTTTCGAT	COCCACCTIT	GTAGTCCGGG	OCAGTOCTOG	TCCCTATCCT	CATOGGANTO	ATAGTGACOO	ACTOCOCION
11101	TAACTACTOC	ט	TTGGGCATTG	ACTTORANGA	CCCATTIAL	ACACANANTO	GAAAACTAGG	ACTANAGEAC	OCCUPANT	TOCATIOTAL .
	ATTICATION		MACCCGTAMC	TOANCITTICE	COCGIANATA	TOTOTITING	CTTTTGATCC	TOATTTICATO	CCCCCAOOAA	ACCTACATIVI
10011			CONTACTAC	TOGITECAGGE	CHCACTATTA	ATAATACTTC	CTTGCMACT	ANGITACTO	GAGCCTTOOO	TTTTGATTCA
77976	The state of the s		GREATEGITS	ACCAGGTCCA	CACTGATAAT	TATTATCAAG	GAACGITIGA	TTTCAATOAC	CTCCOAACCC	MANCTAN .
1		?	4		S. Marie B. State C. St.	PAAPTAGATTG	CTTATACTTO	ATCITACTTA	PCCONTROAT	GCTCANARC:
31301	CANGGCAATA		TUTAGCAGGA	GCAC TANKA	וויאוורורא		Cabrancas	TACAATCAAT	ACCOMACTA	CCACTITION
	Griccortal	ACCITICAATT	ACATCGTCCT	CCTGATTCCT	AACTAAGAGE		פעשואווסיטיב			Charles A.A.
31401	AACTAAATCT	ANGACTAGGA	CAGGGCCCTC	TITITATANA	CTCAGCCCAC		TTAACTACAA	CAMOOCCTT	ALCIDITAL.	
1	TEATTTAGA	TICTOATCCT	GICCCGGGAG	ANANTATIT	GAGTCGGGTG	TTGNACCTAT	AATTGATGTT	CITICCOOM	ATGAACAAAT	GICOMOI II
		Hendill								
11501	CARTICCAAA	AACCTTGAGG	TTAACCTAAG	CACTGCCAAG	COCHICATGE	TTCACCCTAC	AGCCATAGCC	ATTAATOCAG	GAGATOGOCT	TOWATTAGE
1	TITAAGGITT	THEOMACTEC	AATTCGATTC	GTGACGGTTC	CCCAACTACA	AACTOCGATG	TCCCTATCCC	TAATTACGTC	CTCTACCCGA	ACTIMACEA
11601	TCACCTAATO	_	AAATCCCCTC	AAAACINAAA	TYRICCATOR	CCTAGAATTT	GATTCAAACA	AGGCTATGGT	TCCTAAACTA	COMCINATI
1	ACTOCATTAC		TTTAGGGGAG	THOME	AACCCOSTACC	GCATCTTANA	CTAACITICT	TCCCATACCA	ACCALLICAT	רנווושרנות
11701	Transport	2	OCCATTACAG	TAGGNAACAA	MATANTOAT	ANGCTAACTT	TYTHERENCEAC	ACCAGCTCCA	TCTCCTANCT	GTAGACTAGA
	AATCAAAACT	9	COGTANTOTC	Ancomment	TTTATTACTA	TICCATTGA	ACACCTYATE	TOCICOAGGT	ACAGGATIGA	CATCICALIT
11801	TENTECHOLOGY		TCACTITIOGE	CTTAACAAA	TUTAX	MATACTTGC	TACAGITITICA	OFFICER	TTAAAGGCAG	TTTGGCTCCA
1	ACOTOTOTO		AGTCAAACCA	GAATTGTTT	ACACCCTICAG	TTTATGNACO	ATCICANGE	CAAAACCGAC	AATTICCOIC	AAACCOAGO
			TYN TY ATTEN	ATTATAGAT	THENCEANA	TEXAGEGETA	CTANACAATP	CCTTCCTOCA	CCCAGAATAT	T. AACTITA
10616	TATACACCT	ב ב	ACCINCTAGAA	TAATATHETA	ACTOCITIT	ACCTCACGAT	CATTICITION	OGANOGACCT	OCCUCITATA	ACCITICAMAT
	4									
10001	CHARAC	STATE ACTORA	COCACACAC	ATACAMACGE	TGTTCARTT	ATRICCTANCE	TATCACCTTA	TCCMMATCT	CACOCTAAAA	CTGCCMANG
10000		CTTTACCTCT AGNATICACTT	CCCTCTCCCA	PATCHTACG	ACAACCTAAA	TACGGATTOG	ATAGTCGAAT	ACCITITINGA	GTGCCATTIT	GACGGTTTTC

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32101	TAACATTIGIC	AGTCAAGTTT	ACTTANACCG	NEACANANCT	AAACCTTGTAA	CACTAACCAT	TACACTAGAC	GCTACACAGG	NACAGGAGA	CACAACTC!
	ATTICTAACAG	ħ	TGAATTTRACC	TURTHUM	TTTX 3.14CATT	GICATICGTA	ATGTGATETG	CCATGTOTCC	THICHCLE	CTCTTCAC:
32201	AGTOCATACT	CTATOTCATT	TTCATCCCTG	Transchoors Archingges	ACAN TACAT TOTTVIATUTA	TANTGAMATA	TTTCTCACAT NACCATTCTA	CCTCTTACAC GCAGAATGTC	TTTTTCATAC AAAAAGTATO	ATTRECCEAN: TAACGREETS
32301	AATAAAGAAT		ATCITITCAAC	CACAMATAN	TTCAATTITY'A AAGTPAACGT	GAAAATTICA	NATCATTITIT TEMETAAAAA	CATTCAGTAG	TATAGCCCCA	CCACCACATA GGTGGTGTA
32401	GCTTATACAG		CTTANTCAM	CTCACAGAAC	CCTACTATTC	AACCTOCCAC	CHCCCTCCCA	ACACACAGAG Testestest	TACACAGTCC	AAAGAGAG
	COMPTATOL			CALCION CO.	CACOUNTE AND PROPERTY.	O LOW WIND IN	CACOMMEN		A A COCHEANT	ACTION TO THE
32501	CCACCCOMI	TITICOTAGE	TATCATGGGT ATAGTACCCA	AACAGAGATA TTGTCTGTAT	ANTANTCCAC	MINIMET	OTTICCAAAGE	ACAGCTCCCT	TTCCCAGTAG	TCACTATAL
32601	ATAMCTCCC	COOCCAGCTC	ACTTAAGTTC TGAATTCAAG	ATGTCCCTGT	CENTRACTE	ARCCACAGGC	ACGACAGGTT	CTTCCCCTTC	CTTAACGGGC	CCCCTTCCT"
		}					JE;			
32701	AAGTCCACGC	AAGTECACOC CTACATOGO	Gradacteat	AATCGTGCAT	CAGGATAGGG	CONTROTOCT GCAGCAGEGE	GCAGCAGCGC	GCGAATAAAC	TOCHOCCOCC	OCCUCACOCA
	TrcAporoco	DATOTACCCC	CATCTCAGTA	TTACCACGTA	GTCCTATCCC	הככתככתכנא	CGTCGTCGCG	CCCTTAITTO	ACGACGCCOG	CCCCCACCCA
	Pett									-
32801	CCTGCAGGAA	TACAACATOO	CACTOCICIC	CTCAGCGATG	ATTCGCACCG	CCCCGCAGCAT	AAGCGCCTT	GTCCTCCOGG	CACAGCAGCG	CACCETION!
	GGACGICCTT		GTCACCAGAG	GAGTCCCTAC	TAAGCGTTAT	GCCCTCOTA	TTCCCCCCAA	CAGGAGGCCC	Grancercac	GTCCCACT**1
			Part							
12901	TCACTTARA	CAGCACAGTA	ACTGCAGCAC	AGCACCACAA	TATTGITCAN	AATCCCACAG	TGCAACIGCGC	TOTATCCAM	CCTCATOOCO	OCCACCACAG
	AGTGAATTTA		TOACCITCGTG	TCCTOGTGTT	ATAACAAGIT	TTAGGGTGTC	ACGITECTE	ACATAOGTTT	CONGINACOCIC	cccroman
33001	AACCCACGTG	GCCATCATAC	CACAAGCGCA	CCTACATTAA	GTARGERACE	CTCATABACA	CGCTGGACAT	ANACATTACC	TCTTTTGGCA	TOTTGTAATT
	TIGGOTGCAC		GIGTICGCGT	CCATCTAATT	CACCGCTGGG	GAGTATTTGT	CCCACCTGTA	TTTGTAATOG	AGNAMACCOT	ACAACATTAA
		Kprf								
33101	CACCACCTCC	_	TAAACCTCTG	ATTAAACATO	GCGCCATCCA	CCACCATCCT	AMCCAGCTO	GCCAAAACCT	مدددمددمود	TATACACTON.
	GTGGTGGAGG		ATTTOCACAC	TAATTTGTAC	CGCGSTACGT	GOTTOOTAGGA	TITIONITION	COCTITICOS	0000000000	ATATIOTOS:
	Part						•	Enofiv		
10615	ACCOMANCES	CACTOCAACA	ATGACAGITE	AGAGCCCAGG	ACTOSTANCO	ATCICATICATIC	ATCCTCGTCA	TCATATCAAT	OTTOCCACAA	CACAROCACA
	recerronce	CTOACCTTOT	TACTUTCACC	rereagatee	TOACCATTOS	TACCTAGTAG	TACCAGCAGT	ACTATAGTTA	CACCGIOTT	GREFICETET
	-									***************************************
33301	COTOCATACA	CTRCCTCAGG	ATTACAMOCT	CCTCCCCCTT	TACAACCATA	TCCCAGGGAA	CAACCCAFTC	CTGAATCAGC	GIAMATCCCA	CACTGCAGG
	OCACOTATOT	DANDGAGTCC	TANTOTTCGA	CHANDRICOCA	ATCTINGTAT	ACCEPTE	CTTCACTANG	GACTTAOTCO	CATTINGGGT	פופאכנוני.
33401	AAGACCTOC	ACCITANCICA	CGTTGTGCAT	TOTOMOTO	TTACATTCCS	CATACACROTA	ATOATCCTCC	AGTATOOTAG	CCCCCCTTTC	TCTCTCAM
	TTCTOGAGCO	TOCATTOAGE	OCANCALGTA	ACASTETICAC	ANTOTANGE	נישכישכישיב	TACTACASAGO	TCATACCATC	OCCICCOMAG	ACAGAGETTE
33501	CCACCTACAC	GATCCCTACT	GTACGRAFTS	CRECCOAGACA	ACCICAGATOR	REPRESENTE	ACTOPCATE	CANATOGANC	CCCCACOTA	CTCATATTF.
	CCTCCATCTO	CTAGGGATGA	CATACCTICAC	acochenor	TOCCTCTAGE	ACAACCAACA	TCACAGTACG	GTTTACCTTO	COCCTOCAT	Charathasi

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NESCICENTA GATCOCTETO TOTAGTAGIT GTAGTATATE CACTETETA ACCOCCAAT CTANCARC ACATEANCAA CATCATATAG GACGACGGG ACTATTGTAG GYGGTGGGG CTATTTCGGT GYGGGITCON CARCAACGA ACTATTGTAG GYGGTGGGGT CTATTTCGGT GYGGGITCON GAAGAACCAT GTTTTTTTT TTATTCCAAA AGATTATCCA AAACCTI'NAA	AGANCAGATA ATGGCATTIG TCTTGTCTAT TACCGTAAAC ATCTCCTCTA TAAACATTCC TAGAGGAGAT ATTTGTAAGG	CCCGGTANCA TITITAGNES AGGICLOCO TICAANAGAG GAACAITAAC AAAAATACCO AAGATTATGC CTTGTAATTG TITITATGC CTTCCCCCCC AGGAACCATG ACAAAAGAAC GAAGGGGCGG TCCTTGGTAC TGTTTTCTTG	CTRETTGEAT GOGGGGCAT ATANANTOCA AGGTGCTOCT CAAAAATCA GAACAACGTA CCGCCGCTA TATTTTACOT TCCACGACGA GITTTTTAGA ATAAAAACGA GITTTTTAGA ATAAAAAAAAAAAAAAAAAAAAAAAA	TTAMANGCA ANTITITICGT GACCGAAATA CTGGCTTTAT	ACCICION AMECCINE COLIMOCAN FIGURACIA TITOLOGICA COGNICCETT ACTIVADADA AMACCIATA AMANACIA CONTINUI TITOCOMINA TITITITATO CONTINUI CONTINUI TITOCOMINA TITITITATO CONTINUI CONT	ACTOCATIOC CAATITCAGG
ACAGATCTRC CAGAGGCAN CTATGTAANC TCCTTVATGC GATACATTR AGGAAGTACG GAGANGAGCG CGTTCTAGG CCCTCCTCCC CGTTCTAGG	TCCGGTGGC TGGTCAAACT AGGCCAGGC ACCAGTTTGA TCGACGTAAA GGCTAAAAGCC ACCTGCATTT CCGATTTGAG TATCTCTAAG CAAATCCCGA	ATAGAGATTC GTTTAGGGCT AATTCAGGTT CCTCACAGAC TTAAGTCCAA GGAGTGTCTG CGTGCAGGTC TGCACGGAGC GCACGTCCAG ACGTGCCTGG	CAGCGTAGCC CCCATGTAAG GTCGCATCGG GGCTAACATTC TCGTAGTCAT GCTCATGCAG ACCATCAGTA CGAGTACGTC ACACAAAATA AAATAACAAA	GCGTGGCGF ANAMAGTG CCGCCTGGCG TTTTTTTGC GFTGATTCAC ATCGGTGAGT CAACTAAGTG TAGCCAGTCA	ANCHANATA ATAGONGAGA TEGETETANE TATECTETE ACAGGGGA CCATANGAGE TGEGCCGTC GGTATEGTCA	ANDORCCANG TECNFACEGA TTCCCGGTTC ACGTCTCGCT
CANA ACCAGGIGG GEGGTGACAA OTTT TGGTCCACGC CCGCATTTT GGTC GCGCCCCTG ICTTCGGGTT GGTC CCGGGGGC CGAAGCCCAA ACAA TCGTTCTGCG ACTCACCCAA TCGTTCTGCC ACTCACACCCAA	ATGALAGET ATTARGIGAA CGCGCTCCCC TACTTCTAGA TAATTCACTT GCGCGAAAAAAAAAAAAA	TAGAGCGATCA CAGCGATCA GYCCCTTAGT CAGGCCAGC GYCCCGGTCG	ATACTCOGAG ' TATGAGCCTC COCCCAAAAA OCOCCTTTTT GRETGCCGGT	HOTA CAGACOCCA ANGACOTATT ATAA GACOOACTAC GGCCATGCCG FIATT CTGCCTGATG CCGGTACGGC ATOTA AGACTCCGTA AACACATCAG	ACAGCCCCA TOTCGGGGGT GAACAACATA CTIGTIGTAT	CAGCT CAATCAGTCA CAGTGTAAAA OTCGA GTTAGTCAGT GTCACATTTT
13601 CTGMACCAMA GACTTCOTTT 33701 AAGCATCCAG TTCOTAGOTC 53801 ACCTACACAT TGGATGTAA		34201 CAGCCTCANG GACCTCANG GACGGAGGTC J4301 COTCCCTTCG CCAGGGAAGC	34401 TATGACACCC ATACTOTOCC 34501 COCANACCT CCGTTTCCGA	AGAGTTTGTA 34701 ATACCATAA TATTCGTATT 34801 TCATAATGTA	34901 AGACAACAT TCTGTTGTAA 35001 TCCCGCTCCA AGACCAAGGT	35101 GOCACCAGCT CCGTGOTCGA

Figure 15V

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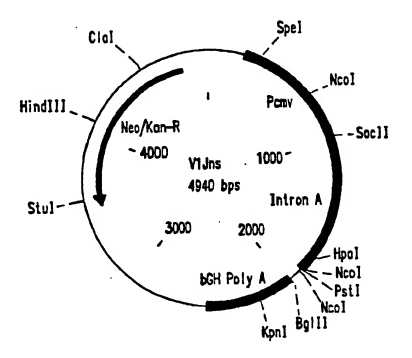
35301	CATTITAAGA	AAACTACAAT	TCCCAACACA	TACAAGTTAC	TrecheceTAA Magazaggata	AACCTACGIC	Acendetect	TTCCCACGC	CCCCCCCACO	TCACAMETE
						Pa	75			
•	•						Fcoffi			
35401	CACCCCTCA		GCCTTCAATC	CANATANCE	TATATTA	ATCATCTTAA	TTANGAATIC	GGATCTGCGA	COCCAGGCTO	GATGGCCTT.
	GTOGGGGAGT	AATAGTATAA	CCGAAGTTAG	GTTTTATTCC	ATATAATAAC	TACTACAATT	AATTCTTAAG	CCTAGACGCT	OCOCIACONE	בדאכנינסטאט
35501	CCCATTATOA		Treededec	ATCGGGATGC	CCGCGTTGCA	COCCATOCTO	TCCAGGCAGG	TAGATCIACGA	CCATCAGGGA	CAGCTTCAAG
	CCCTAATACT	AMGRAGAGCG	AMGGCCGCCG	TAGCCCTACG	GGCGCAACT	ככנציואכפאכ	AGGICCOICC	AILIALIGET	WOTABILLE.	CICOMO II.
35601	GCCAGCAAAA		CCTTAAAAAQCI	CCCCOTTGCT	GGCGTTTTTTC	CATAGRETEC	GCCCCCCTOA	CONGCATCAC	ANAMATICARC	GCTCAAGTC/.
	coorcornin	CCCCTCCTTG	CCATTITICE	GOCGCNACGA	CCCCAAAAG	GTATCCCAGG	CCCCCCCCACT	OCTCGTAG10	TTITTAGCTO	CGAGTTCAGF
35701	CACCTCCCCA		_	ATACCAGGCO	THECCECTO	GAAGCTCCT	CONCOCICT	CCTOTICCOA	CCCTOCCGCT	TACCCCATAC
	CTCCACCGCT	TIGOGCIOIC	CTGATATITC	TATOSTCCOC	AAAGTGGGAC	CHICOAGOCA	GCACCCCAGA	CCACAAGCCT	CKITACGGCGA	ATGGCCTATE
35801	CTUTCCOCCT		OCCANGCISTS	CACCE TITLE	ATARCTCACG	CTCTACKGTAT	CTCANTTICOG	TOTAGGTCOF	Tractccand	כיומסמכיומדי.
	GACAGGCGGA	AAGAGGGAAG	CCCTTCGCAC	COCCANACAG	TATCGAGTOC	GACATCCATA	GARTICANCIC	ACATCCAGCA	AGCGAGGTTC	DACCCGACA!
35901	TOCACOAACC	CCCCOPICAG	CCCGACCGCT	GCGCCTTATC	CCCTAACTAT	CONCINGAGE	CCAACCCGGF	AAGACACGAC	TTATCOCCAC	TOOCAGCAGY.
	ACCITOCITION	GOCCAAGTC	OCCUPACION	COCOGNATAG	GCCATTGATA	GCAGAACTICA	GCITICOCCCA	ricidideno	AATAGCOOTG	ACCGROCTIC
36001	CACTGGTAAC	AGGATTAGCA	GACCGACGTA	TOTAGGCGGT	GCTACAGAGT	TETTRANSTO	GTOCCCTAAC	TACOGCTACA	CTAGANGGAC	AGTATTICOF
	STURCCATTO		CTCGCTCCAT	ACATECGCCA	CGATGICTCA	AGNACTITCAC	CACCOGATTO	ATOCCOATOT	DATICTACCIO	TCATAAACCA
36101	ATCTOCGCTC	TOCTONAGEC	AGTTACCTTC	CCANANACAG	TICKSTANCIC	THEATCEGGE	ANACARACCA	CCCCTGGTAG	COORGOTTT	TTTGFTTVE.
	TAGACGCGAG	ACCACTTCGG	fcantocand	CCTTTTTCTC	NACCATCGAG	AACTAGGCCG	THEFTIGGE	GOCGACCATC	OCCACCAAAA	AACAAACCT
36201	AGENGCAGAT	TACGCGCAGA	AAANANGGAT	CTCAAGAAGA	TCCTTTGATC	TITICTACOO	CONCINCACCIC	TCAGTOGNAC	GAMACTCAC	GTTMOCCA!
	ACCICCICTA	ATGCGCGTCT	TETTTTCCTA	GAGITCTICT	ACCIMACTAG	MANGATOCC	CCAGACTACO	AGREACETTO	CHITICAGIO	CAATTCCCTA
36301	THOOTCATO	AGATTATCAA	AAAGGATCTT	CACCTAGATO	CITTIAMATIC	ANTCHANGE	ATATATOAGT	AAACTTKGGTC	TOACACTTAC	CNATCCTEN
	AAACCAGTAC		THICCTAGAA	GTGGATCTAG	GAAAATTTAG	TTAGATTTICA	TATATACTCA	THICANCCAG	ACTOTICAATO	CTINCOLATI
36401	TCAGTGAGGC	ACCTATCTCA	OCGATCTOTC	TATTACGITE	ATCCATAGIT	OCCITOACTOC	CCGTCGTGTA	GATAACTACG	ATACOGGAGG	GCTTACCATY.
	AGTCACTCCO	TOGATAGAGT	CGCTAGACAG	ATARAGEME	TACCTATCAA	CCACACACACO	GGCAGCACAT	CTATHCATCC	TATOCCCTCC	CGNATCGTAG
36501	TOOCCCCAOT	GCTGCAATGA	TACCCCCAGA	CCCACOCTCA	CCGCCTCCAG	ATTTATCAGE	AATAAACCAG	CCACCCCCAA	GOGCCGAGCG	CACIANGRAGE
	ACCOGGGTCA	CGACGITIACT	ATGGCGCTCT	OCCIOCOAST	CCCCCACGIC	TAAATAGTCO	Trattrogre	gorcoccTT	CCCOOCICOC	CHCTTCACCA
36601	CCTGCAACTT	FATCCOCCTC	CATCCAOTCT	ATTAATTCIT	GCCGGGAAGC	TAGASTAAGT	AGITCGCCAG	TTAATAGTIT	OCOCAACGTT	GITTCCCATTG
•	OCACOTIONA		GTAGGTCAGA	TAATTAACAA	COCCCTTCG	ATCTCATTCA	TCANGCOUTC	AATTATCAAA	COCOTTOCAA	CANCOGTAAC
36701	CTACAGGCAT		COCTCGTCGT	TTOOTATOOC	TTCATTCAGC	TCCGGTTCCC	AACGATCAAG	GCCAOTTACA	TCATCCCCCA	TOTTOTACAA
	GATOTCCOTA		GCGAGCAGTA	AACCATACCG	AAGTAAGTCO	AGCCANGGG	TIGGTAGTIC	COCTCAATOT	ACTAGGGGGT	ACAACACGTT
			<u> </u>	_*						
36801	AAAAGCOGTT	ACCICCTICO	GICCICCOAT	Greenecoar contentals	ACTAACTICG	CCGCAGTGTT	ATTENCHENTS	CITATOCCAG	CACTGCATAA	TICTCTTACT
	TTTTCGCCAA		CAGGAGGCTA	GCAACAGTICT	TCATTCAACC	GOCCITCACAA	TACTICACTAC	CANTACCOTC	GTGACGTATT	ANGAGAATUA
36901	GTCATGCCAT	CCGT	cerrector		ACTUACCAA		CAATACHCTA	MOCHOCOACC	CHCAACGAGA	ACCCC(ASCC)
	CAGTACGGTA	COCATTICTAC	GAMAGACAC	TUNCCACTOR	Tener I tener	ראניומטירו				

figure 15W

PMRKAd5qag MER682

TACCCECTETT	TCTCTCATCA ACAGAGTAC"	CCATTATTA': GCTAATAATA	
CANCACGODA FANTACCOCO CCACATARCA GAACTITAAA AGTRETEATE ATTARAAAAC GITCITGOG GCGAAAGTE TCAAGGATCT TACRICIGITA GATTOCCCC ATTATOCCC COCTITIOAG AGTRECTAGA ATGREGAAAA GAGAACCAT ATTATOCCC COCTITIOAG AGTRECTAGA ATGREGAAAA GAGAACCAGT TCAACTAGAACCAGA ACCAACTAGA GATTOCCC ACCACACTAGA ATGRAAAA CACAACTAGA GATTOCCC ACCACACTAGA AGCAACAAAA GACCACATAGA GATTATOCCC ACCAACTAGA AGCAACAATAGA AGCAACAATAGA AGCAACAATAGA AGCAACAATAGA AGCAACAATAGA AGCAACAATAGA AGCAACAAAA AGCAACAATAGA AGCAACAAAAAAAAAA	OCCOCANAA AGGINTAAG OCCACACAS AAATCITISAA TAFICATAGI FITCCITITI CAATAITAT GAAGCAFITA TCAGGITAT TCTCITAAA. CGGCOFFITI TCCCTIATIC CCCCRICKSC TIFACAACIT ATSAGIAATA GAAGAAAAA GITAAAAAA CFICOTAAAF AGTCCCAATA ACAGAGTACI'	CCCCANGRA ATTICANT ATTICANA ATANCANA AGGITTICE CENTANTE CCCCANAGE CCCTATACA CAGGACTE CAGANTATT CATANTANA CCCTANGRA INANCITACA TANCITACA TANTETTT TATTICITA TCCCCANAGE CCCTANAGE) ID NO: 27) ID NO: 28)
STATTERAGE CONTRACTERAGE CONTR	NGCATTT!	CACCTGAC	MT (SEC
33 53 53 53 54 45 54 45	35 X	10 40 V	CT TA
GTTCTTCG CAAGAAGG CACCAGG	CAATATTA	CCCGAAAAA GGGCTTTTC Feelil	Bamfil GGA TUCGAATT
ATTCS:AAAAC TAACCTTTTG CTTTTAC:TFT GAAAATAAAA	CHCCTTTTT	CCTCTANAGE GCCTCTANAGE	BA NAGANTINGA THETTANGET
AGTOS, TEATO TY, AGGAGTAG TY TY, AGA ATTATA	TACTCATACT ATGACTATEA	AGGGS STITCES TECECEAAGGE	TTICGTCTTC AAAGCAGAAG
GANCTITAAA CITGAAATIT ACCCAACIGA TAAGITGACI	AAAATCITTGAA TTFACAACTT	ATAMACAAAT TATTTGTTTA	CACGAGGECE
CCACATAGGA GGTGTATCGT CCACTCGTGC GGTGAGGCACG	OCCGNSTOCC CCGCWTGCC	ATTTAGAAAA TAAATCTTTT	ATACECCATA TATCCCCATA
TANTACCOCO ATTATOCCOC TCOATGTAAC	AGGINATAG	ATTTGAATGT TAAACTTACA	CATOACATTA ACCTATAAAA ATACACGTAT CACGACACCC TTICGTCTTC AAGAATTCGA TCGAATTCT TAAT (SEQ ID NO: 27) GIACTOTAAT TGATATTT TATCCGCATA GTGCTCCAGG AAAGCAGAAG TTCTTAACCT AGGCTTAAGA ATTA (SEQ ID NO: 28)
	CCCCANAN	CCCCTATOTA	
37001	37201	37301	37401

Figure 15X



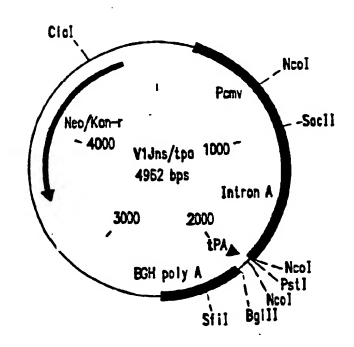


FIGURE 16

GCAGTGGCCCCTGACTGAGGAGAAGATCAAGGCCCTGCTGGAAATCTGCACTGAGATGGAGAAGGAGGGCAAAATCTCCA sGInTrpProLeuThrGiuGiu_yslieLysAioLeuVoiGiuIieCysThrGiuMelGiuLysGiuGiyLysIieSerL 30 40 50

AGATIGGCCCCGAGAACCCCTACAACACCCCTGTGTTTGCCATCAAGAAGAAGAAGAACCACTCCACCAAGTGGAGGAAGCTGGTG
ys!!eG!yProG!uAsnProTyrAsnThrProVo!PheA!o!!eLysLysLysAspSerThrLysTrpArgLysLeuVo!
60 70

GACTICAGGGGGCTGAACAAGAGGACCCAGGACTTCTGGGAGGTGCAGCTGGGCATCCCCCACCCCGCTGGCCTGAAGAA AspPheArgGluLeuAsnLysArgThrGlnAspPheTrpGluVolGlnLeuGlylleProHisProAloGlyLeuLysLy 80 90 100

GAAGAAGTCTGTGACTGTGCTGGCTGTGCCGGATGCCTACTTCTCTGTGCCCCTGGATGAGGACTTCAGGAAGTACACTG slyslysSerVolThrVolLeu<u>Alo</u>VolGlyAspAloTyrPheSerVolProLeuAspGluAspPheArgLysTyrThrA 110 120 130

CCTTCACCATCCCCTCCATCAACAATGAGACCCCTGGCATCAGGTACCAGTACAATGTGCTGCCCCAGGGCTGGAAGGGC InPhetnrlieProSerlieAsnAsnGluthrProGly!leArgTyrGinTyrAsnVolLeuProGlnGlyTrpLysGly 140 150

TCCCCTGCCATCTTCCACTCCTCCATGACCAAGATCCTGGAGCCCTTCAGGAAGCAGAACCCTGACATIGTGATCTACCA SerProAlollePheGinSerSerMetThrLysTieLeuGluProPheArgLysGinAsnProAsplieVollleTyrG1 160 170 180

TGCTGAGGTGGGGCCTGACCACCCTGACAAGAAGCACCAGAAGGAGCCCCCCTTCCTGTGGATGGGCTATGAGCTGCAC euleuArgTrpGlyLeuThrThrProAsplysLysHisGInLysGIuProProPheleuTrpMetGlyTyrGIuLeuHis 220 230

CCCGACAGTGGACTGTGCAGCCCATTGTGCTGCCTGAGAAGGACTCCTGGACTGTGAATGACATCCAGAAGCTGGTGGG ProAsplysTrpThrVoIGinProlieVoiLeuProGluLysAspSerTrpThrVoiAsnAsplieGinLysLeuVoIGI 240 250 260

CAAGCTGAACTGGGCCTCCCAAATCTACCCTGGCATCAAGGTGAGGCAGCTGTGCAAGCTGCTGAGGGCCACCAAGGCCC yLysLeuAsnTrpAloSerGInlleTyrProGiy11eLysVolArgGInLeuCysLysLeuLeuArgGiyThrLysAloL 270 280 290

FIGURE 17A

GCCCTGTACTATGACCCCTCCAAGGACCTGATTGCTGAGATCCAGAAGCAGGGCCAGGGCCAGTGGACCTACCAAATCTA G1yVo1TyrTyrAspProSerLysAspLeu1teAloGiu1teGlnLysGInGlyGInGlyGInTrpThrTyrGin1teTy 320 330 340

CCAGGAGCCCTTCAAGAACCTGAAGACTGGCAAGTATGCCAGGATGAGGGGGGGCCCACACCAATGATGTGAAGCAGCTGA rGInGIuProPheLysAsnLeuLysThrGIyLysTyrAloArgMelArgGlyAloHisThrAsnAspVoiLysGInLeuT 350 350 370

CTC4GGCTGTGCAGAAGATCACCACTGAGTCCATTGTGATCTGGGGCAAGACCCCCAAGTTCAAGCTGCCCATCCAGAAG hrGluAloVolGinLysIleThrThrGluSerlleVollleTrpGlyLysThrProLysPheLysLeuProlleGinLys 380 390

GGTGAAGCTGTGGTACCAGCTGGAGAAGGAGCCCATTGTGGGGGGCTGAGACCTTCTATGTGGCTGGGGCTGCCAACAGGG uVollysleuTrpTyrGinleuGiuLysGiuProlleVolGlyAloGiuThrPheTyrVolAloGlyAloAloAsnArgG 430 440 450

AAGACTGCCCTCCAGGCCATCTACCTGGCCCTCCAGGACTCTGGCCTGGAGGTGAACATTGTGACTGCCTCCCAGTATGC
LysThrAioLeuGinAiolieTyrLeuAioLeuGinAspSerGiyLeuGiuVoiAsnIieVoiThrAioSerGinTyrAi
480
490
500

CCTCGGCATCATCCAGGCCCAGCCTGATCAGTCTGAGTCTGAGCTGGTGAACCAGATCATTGAGCAGCTGATCAAGAAGG oleuGiyiieiieGinAioGinProAspGinSerGiuSerGiuLeuVolAsnGinIieiieGiuGinLeuIieLysLysG 510 520 530

ACAACGTGTACCTGCCTGCCTGCCCACAACGCCATTGGGGGCAATGAGCAGGTGGACAAGCTGGTGTCTGCTGGC !ulysVo!TyrleuA!oTrpVo!ProA!oHislysG!y!!eG!yG!yAsnG!uG!nVo!AsplysLeuVo!SerA!oG!y 540 550

ATCAGGAAGGTGCTGTTCCTGGATGGCATTGACAAGGCCCCAGGATGAGCATGAGAAGTACCACTCCAACTGGAGGGCTAT
11eArgLysVolleuPheleuAspGly11eAspLysAloGInAspGluHisGluLysTyrHisSerAsnTrpArgAloMe
560 570 580

FIGURE 17B

GGCCTCTGACTTCAACCTGCCCCCTGTGGTGGCTAAGGAGATTGTGGCCTCCTGTGACAAGTGCCAGCTGAAGGGGGAGG taloSerAspPheAsnLeuProProVolVolAloLysGluIleVolAloSerCysAspLysCysGlnLeuLysGlyGluA 590 600 610

GCTGTGCATGTGGCCTCCGGCTACATTGAGGCTGAGGTGATCCCTGCTGAGACAGGCCAGGAGACTGCCTACTTCCTGCT AlovalHisvalAlaSerGlyTyrlleGluAlaGluVallleProAlaGluThrGlyClnGluThrAlaTyrPheLeuLe 640 650 660

GAAGCTGGCTGGCAGGTGGCCTGTGAAGACCATCCACACTGCCAATGGCTCCAACTTCACTGGGGCCACAGTGAGGGCTG

uLysleuAloGlyArgTrpProVolLysThrIleHisThrAloAsnGlySerAsnPheThrGlyAloThrVolArgAloA

670

680

CCTGCTGGTGGGCTGGCATCAAGCAGGAGTTTGGCATCCCCTACAACCCCCAGTCCCAGGGGGTGGTGGCCTCCATGAAC
IoCysTrpTrpAioGly!ieLysGinGluPheGly!ieProTyrAsnProGinSerGinGlyVolVolAIoSerMelAsn
700 710

AAGGAGCTGAAGAAGATCATTGGGCAGGTGAGGGACCAGGCTGAGCACCTGAAGACAGCTGTGCAGATGGCTGTGTTCAT LysG1uLeuLysLys11e11eG1yG1nVo1ArgAspG1nAloG1uHisLeuLysThrAloVo1G1nMetAloVo1Phe11 720 730 740

CCACAACTTCAAGAGGAAGGGGGCATCGGGGGCTACTCCGCTGGGGAGAGGATTGTGGACATCATTGCCACAGACATCC
eHisAsnPheLysArglysGlyGlylleGlyGlyTyrSerAloGlyGluArglleVolAsplleIleAloThrAsplleG
750
770

AGACCAAGGAGCTCCAGAAGCAGATCACCAAGATCCAGAACTTCAGGGTGTACTACAGGGACTCCAGGAACCCCCTGTGG
InThrLysGTuLeuGInLysGInTleThrLysTleGInAsnPheArgVoITyrTyrArgAspSerArgAsnProLeuTrp
780 790

AAGGCCCTGCCAAGCTGCTGTGGAAGGGGGAGGGGGCTGTGGTGATCCAGGACAACTCTGACATCAAGGTGGTGCCCAG LysGtyProAtoLysLeuLeuTrpLysGtyGtuGtyAtoVotVotIteGtnAspAsnSerAspIteLysVotVotProAr 800 820

AAAGCCCCGGCAGATCT (SEQ ID NO: 3)
Xx Bq 11 (SEQ ID NO: 4)

FIGURE 17C

RoSerGIUIIeSerAloProlieSerProlieGiuThrVoIProVoiLysLeuLysProGlyMelAspGly 20 20

FIGURE 18

WT	- ATG GGT GGC AAG TGG TCA AAA CGT AGT GTG CCT GGA TGG TCT	-42
OPT	- ATG GGC GGC AAG TGG TCC AAG AGS TCC GTG CCC GGC TGG TCC M G G K W S K R S V P G W S	-14
WT	- ACT GTA AGG GAA AGA ATG AGA CGA GCT GAG CCA GCA GCA GAT	-84
OPT	- ACC GTG ÁGG GÁG ÁTG ÁGG ÁGG GCC GÁC GCC GCC GÁC T V R E R M R R A E P A A D	-28
WT.	- AGG GTG AGA CGA ACT GAG CCA GCA GCA GTA GGG GTG GGA GCA	-126
OPT	. AGG GTG AGG AGG ACC GAG CCC GCC GCC GTG GGC GCC R V R R T E P A A V G V G A	-42
WT	- GTA TCT CGA GAC CTG GAA AAA CAT GGA GCA ATC ACA AGT AGC	-168
OPT	. GTG TCC AGG GAC CTG GAG AAG CAC GGC GCC ATC ACC TCC TCC V S R D L E K H G A I T S S	-56
WT	- AAT ACA GCA GCT ACC AAT GCT GAT TGT GCC TGG CTA GAA GCA	-210
OPT	- AAC ACC GCC GCC ACC AAC GCC GAC TGC GCC TGG CTG GAG GCC N T A A T N A D C A W L E A	-7 0.
WT .	- CAA GAG GAT GAG GAA GTG GGT TTT CCA GTC AGA CCT CAG GTA	·252
OPT	- CAG GAG GAC GAG GAG GTG GGC TTC CCC GTG AGG CCC CAG GTG	-84
МТ	- CCT TTA AGA CCA ATG ACT TAC AAG GGA GCT GTA GAT CTT AGC	-294
OPT	- CCC CTG AGG CCC ATG ACC TAC AAG GGC GCC GTG GAC CTG TCC P L R P M T Y K G A V D L S	-98
WT	- CAC TIT TTA AAA GAA AAG GGG GGA CTG GAA GGG CTA ATT CAC	-336
OPT	- CAC TTC CTG AAG GAG AAG GGC CTG GAG GGC CTG ATC CAC H F L K E K G G L E G L I H	-112
WT	- TCA CAG AAA AGA CAA GAT ATC CTT GAT CTG TGG GTC TAC CAC	-378
OPT	- TCC CAG AAG AGG CAG GAC ATC CTG GAC CTG TGG GTG TAC CAC	-126
₩T	- ACA CAA GGC TAC TTC CCT GAT TGG CAG AAC TAC ACA CCA GGG	-420
OPT	- ACC CAG GGC TAC TTC CCC GAC TGG CAG AAC TAC ACC CCC GGC T Q G Y F P D W Q N Y T P G	-140

FIGURE 19A

WT	- CCA GGA ATC AGA TTT CCA TTG ACC TTT GGA TGG TGC TTC AAG -46	2
OPT	- CCC GGC ATC AGG TTC CCC CTG ACC TTC GGC TGG TGC TTC AAG P G I R F P L T F G W C F K -15	4
₩T	- CTA GTA CCA GTT GAG CCA GAA AAG GTA GAA GAG GCC AAT GAA -50)4
OPT	- CTG GTG CCC GTG GAG CCC GAG AAG GTG GAG GCC AAC GAG L V P V E P E K V E E A N E -16	58
WT	- GGA GAG AAC AAC TGC TTG TTA CAC CCT ATG AGC CAG CAT GGG -54	16
OPT	- GGC GAG AAC AAC TGC CTG CTG CAC CCC ATG TCC CAG CAC GGC G E N N C L L H P M S Q H G -18	B2
WT	- ATA GAG GAC CCG GAG AAG GAA GTG TTA GAG TGG AGG TTT GAC -51	88
OPT	- ATC GAG GAC CCC GAG AAG GAG GTG CTG GAG TGG AGG TTC GAC I E D P E K E V L E W R F D -1	96
WT .	ADD THE DIA GOT THE COLOR	30
OPT	- TCC AAG CTG GCC TTC CAC CAC GTG GCC AGG GAG CTG CAC CCC	210
WT	- GAG TAC TAC AAG GAC TGC TGA (SEQ ID NO:30) -6	551
OPT	- GAG TAC TAC AAG GAC TGC TAA (contained within SEQIDNO:9) E Y Y K D C (SEQIDNO:10) -2	216

FIGURE 19B

VIJNSTITET PSEI CATGGGTCTTTTCIGGGGCCTCCTTGAGAICIGCCACC ATG GGC GGC ANG TGG TCC ANG TCC GTG CCC VIJns/nef

SrfI BallI CAC CCC GAG TAC TAC AAG GAC TGC TAA AGCCCGGGGAGAICIGCTGGCCTTCTAGTTGCCAGC (SEQ 1D NO: 38) H P E Y Y K D C * (contained withir SEQ 1D NO: 10:

V1Jns/nef(G2A.LLAA)

Psti Catrosetetttictocaccotecttoagatetoccacc atg gcc ggc ang tgg tec ang agg tec gtg eec

Srff Bollf

CAC CCC GAG TAC TAC AAG GAC TGC TAA AGCCCGGGGAGAICIGCTGCCTTCTAGTGCCAGC (SEQ ID NO: 39)

H P E Y Y K D C * (contained within SEQ ID NO:14)

VlJns/tpanef & VlJns/tpanef(LLAA)

CATEGETCTTTICIECAGETCACCETTATATCTAGATCACC ATG GAT GCA ATG ANG AGA GGG CTC TGC TGT GTG V

CTG CTG CTG TGT GGA GCA GTC TTC GTT TCG CCC AGC GAG Λ IC TCC TCC AAG AGG TCC GTG CCC \dots

SrfI Bg111
.... CAC CCC GAG TAC TAC AAG GAC TGC TAA AGCCCGGGAGAICIGCTGTGCCTTCTAGTTGCCAGC (SEQ 1D ND: 40)
H P. , E Y Y K D C * (contained withon SEQ ID NO: 16)

TGURE 20

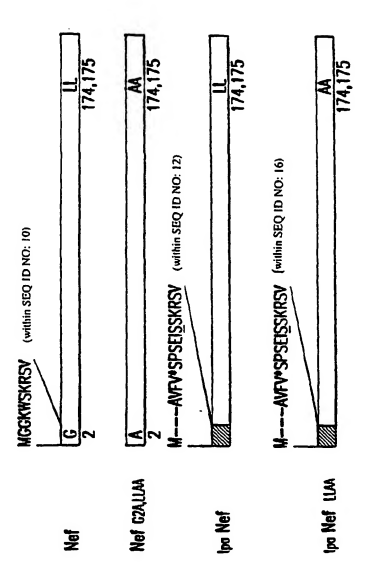


FIGURE 21

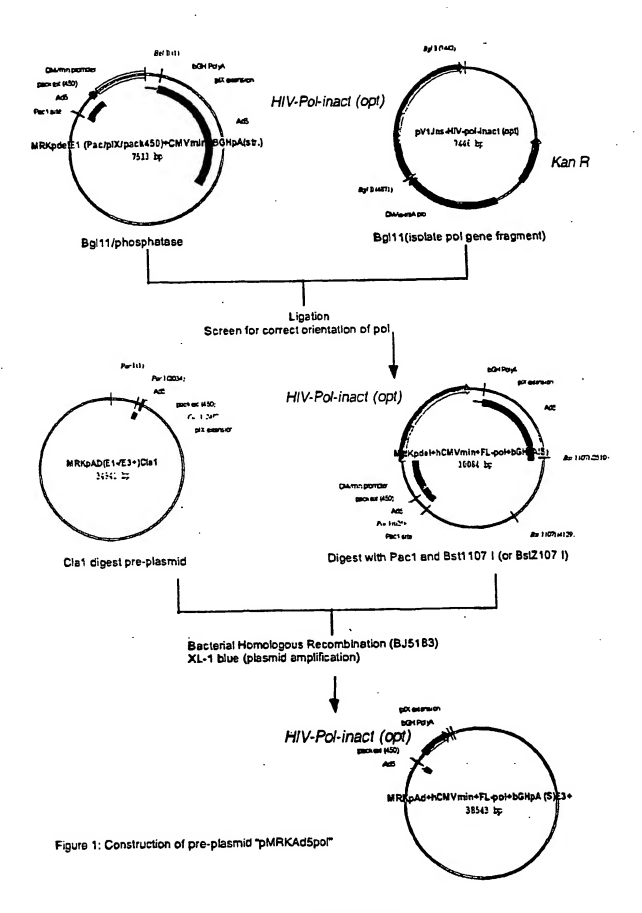
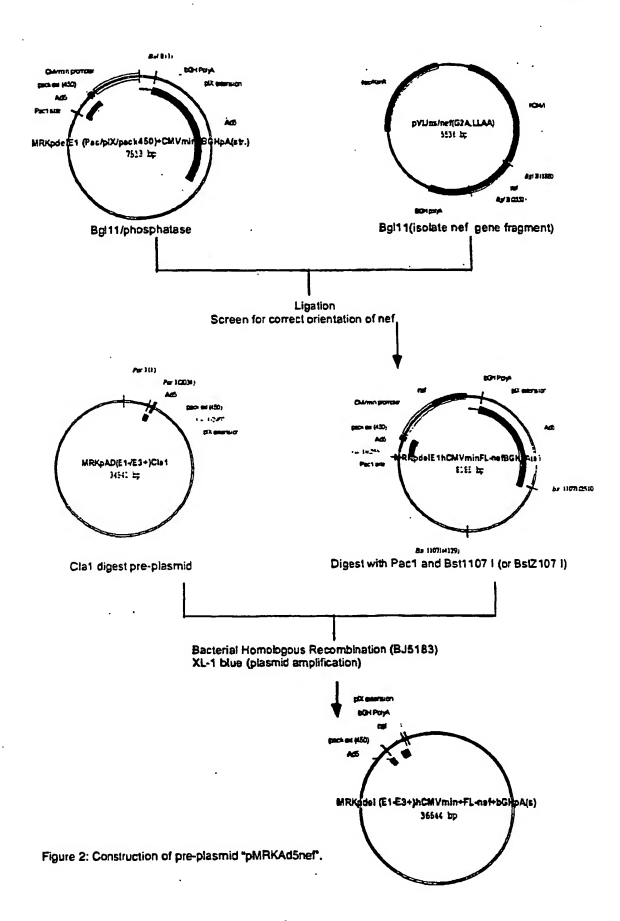
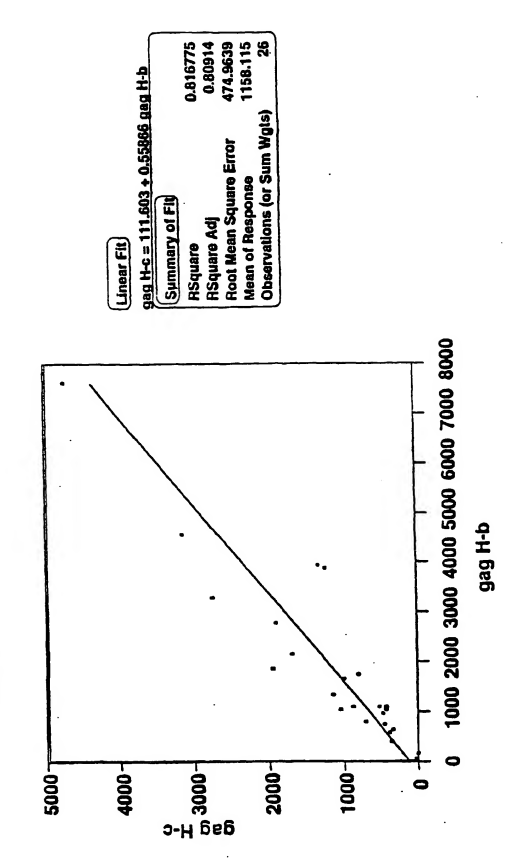


FIGURE 22



Comparison of Clade B vs. Clade C Anti-gag T Cell Responses in Clade B HIV-Infected Subjects



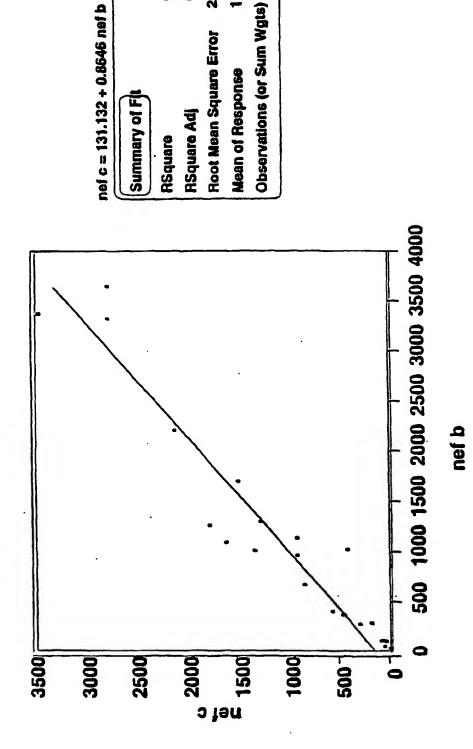
0.91685

289.7718 1096.435

FIGURE 25

ន

Comparison of Clade B vs. Clade C Anti-nef T Cell Responses in Clade B HIV-Infected Subjects



MRKAd5pol MER1062 (MRKAd5 Pre-Adenoviral Vector Containing the IA opt pol Coding Region)

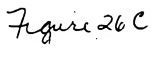
1 CATCATCAAT AATATACCTT ATTTTGGATT GAAGCCAATA TGATAATGAG GTAGTAGTTA TTATATGGAA TAAAACCTAA CTTCGGTTAT ACTATTACTC 51 GGGGTGGAGT TTGTGACGTG GCGCGGGGCG TGGGAACGGG GCGGGTGACG CCCCACTCA AACACTGCAC CGCGCCCCGC ACCCTTGCCC CGCCCACTGC 101 TAGTAGTGTG GCGGAAGTGT GATGTTGCAA GTGTGGCGGA ACACATGTAA ATCATCACAC CGCCTTCACA CTACAACGTT CACACCGCCT TGTGTACATT 151 GCGACGGATG TGGCAAAAGT GACGTTTTTG GTGTGCGCCG GTGTACACAG CGCTGCCTAC ACCGTTTTCA CTGCAAAAAC CACACGCGGC CACATGTGTC 201 GAAGTGACAA TTTTCGCGCG GTTTTAGGCG GATGTTGTAG TAAATTTGGG CTTCACTGTT AAAAGCGCGC CAAAATCCGC CTACAACATC ATTTAAACCC 251 CGTAACCGAG TAAGATTTGG CCATTTTCGC GGGAAAACTG AATAAGAGGA GCATTGGCTC ATTCTAAACC GGTAAAAGCG CCCTTTTGAC TTATTCTCCT 301 AGTGAAATCT GAATAATTTT GTGTTACTCA TAGCGCGTAA TATTTGTCTA TCACTTTAGA CTTATTAAAA CACAATGAGT ATCGCGCATT ATAAACAGAT 351 GGGCCGCGGG GACTTTGACC GTTTACGTGG AGACTCGCCC AGGTGTTTTT CCCGGCGCCC CTGAAACTGG CAAATGCACC TCTGAGCGGG TCCACAAAAA 401 CTCAGGTGTT TTCCGCGTTC CGGGTCAAAG TTGGCGTTTT ATTATTATAG GAGTCCACAA AAGGCGCAAG GCCCAGTTTC AACCGCAAAA TAATAATATC 451 GCGGCCGCGA TCCATTGCAT ACGTTGTATC CATATCATAA TATGTACATT CGCCGGCGCT AGGTAACGTA TGCAACATAG GTATAGTATT ATACATGTAA 501 TATATTGGCT CATGTCCAAC ATTACCGCCA TGTTGACATT GATTATTGAC ATATAACCGA GTACAGGTTG TAATGGCGGT ACAACTGTAA CTAATAACTG 551 TAGTTATTAA TAGTAATCAA TTACGGGGTC ATTAGTTCAT AGCCCATATA ATCAATAATT ATCATTAGTT AATGCCCCAG TAATCAAGTA TCGGGTATAT 601 TGGAGTTCCG CGTTACATAA CTTACGGTAA ATGGCCCGCC TGGCTGACCG ACCTCAAGGC GCAATGTATT GAATGCCATT TACCGGGCGG ACCGACTGGC 651 CCCAACGACC CCCGCCCATT GACGTCAATA ATGACGTATG TTCCCATAGT GGGTTGCTGG GGGCGGGTAA CTGCAGTTAT TACTGCATAC AAGGGTATCA 701 AACGCCAATA GGGACTTTCC ATTGACGTCA ATGGGTGGAG TATTTACGGT TTGCGGTTAT CCCTGAAAGG TAACTGCAGT TACCCACCTC ATAAATGCCA 751 AAACTGCCCA CTTGGCAGTA CATCAAGTGT ATCATATGCC AAGTACGCCC TTTGACGGGT GAACCGTCAT GTAGTTCACA TAGTATACGG TTCATGCGGG 801 CCTATTGACG TCAATGACGG TAAATGGCCC GCCTGGCATT ATGCCCAGTA GGATAACTGC AGTTACTGCC ATTTACCGGG CGGACCGTAA TACGGGTCAT 851 CATGACCTTA TGGGACTITC CTACTTGGCA GTACATCTAC GTATTAGTCA GTACTGGAAT ACCCTGAAAG GATGAACCGT CATGTAGATG CATAATCAGT

7 i jure 26A

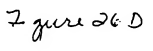
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	AGCGATAATG	GTACCACTAC	GCCAAAACCG	TCATGTAGTT	ACCCGCACCT
951	TAGCGGTTTG	ACTCACGGGG	ATTTCCAAGT	CTCCACCCCA	TTGACGTCAA
		TGAGTGCCCC			
1001	TGGGAGTTTG	TTTTGGCACC	AAAATCAACG	GGACTTTCCA	AAATGTCGTA
		AAAACCGTGG			
1051	ACAACTCCGC	CCCATTGACG	CAAATGGGCG	GTAGGCGTGT	ACGGTGGGAG
		GGGTAACTGC			
1101	GTCTATATAA	GCAGAGCTCG	TTTAGTGAAC	CGTCAGATCG	CCTGGAGACG
		CGTCTCGAGC			
1151	CCATCCACGC	TGTTTTGACC	TCCATAGAAG	ACACCGGGAC	CGATCCAGCC
		ACAAAACTGG			
1201	TCCGCGGCCG	GGAACGGTGC	ATTGGAACGC	GGATTCCCCG	TGCCAAGAGT
		CCTTGCCACG			
1251	GAGATCTACC	ATGGCCCCCA	TCTCCCCCAT	TGAGACTGTG	CCTGTGAAGC
		TACCGGGGGT			
1301	TGAAGCCTGG	CATGGATGGC	CCCAAGGTGA	AGCAGTGGCC	CCTGACTGAG
		GTACCTACCG			_
1351	GAGAAGATCA	AGGCCCTGGT	GGAAATCTGC	ACTGAGATGG	AGAAGGAGGG
		TCCGGGACCA			
1401	CAAAATCTCC	AAGATTGGCC	CCGAGAACCC	CTACAACACC	CCTGTGTTTG
		TTCTAACCGG			
1451	CCATCAAGAA	GAAGGACTCC	ACCAAGTGGA	GGAAGCTGGT	GGACTTCAGG
	GGTAGTTCTT	CTTCCTGAGG	TGGTTCACCT	CCTTCGACCA	CCTGAAGTCC
1501	GAGCTGAACA	AGAGGACCCA	GGACTTCTGG	GAGGTGCAGC	TGGGCATCCC
	CTCGACTTGT	TCTCCTGGGT	CCTGAAGACC	CTCCACGTCG	ACCCGTAGGG
1551	CCACCCCGCT	GGCCTGAAGA	AGAAGAAGTC	TGTGACTGTG	CTGGCTGTGG
	GGTGGGGCGA	CCGGACTTCT	TCTTCTTCAG	ACACTGACAC	GACCGACACC
1601	GGGATGCCTA	CTTCTCTGTG	CCCCTGGATG	AGGACTTCAG	GAAGTACACT
	CCCTACGGAT	GAAGAGACAC	GGGGACCTAC	TCCTGAAGTC	CTTCATGTGA
1651	GCCTTCACCA	TCCCCTCCAT	CAACAATGAG	ACCCCTGGCA	TCAGGTACCA
	CGGAAGTGGT	AGGGGAGGTA	GTTGTTACTC	TGGGGACCGT	AGTCCATGGT
1701	GTACAATGTG	CTGCCCCAGG	GCTGGAAGGG	CTCCCCTGCC	ATCTTCCAGT
	CATGTTACAC	GACGGGGTCC	CGACCTTCCC	GAGGGGACGG	TAGAAGGTCA
1751	CCTCCATGAC	CAAGATCCTG	GAGCCCTTCA	GGAAGCAGAA	CCCTGACATT
•		GTTCTAGGAC			
1801	GTGATCTACC	AGTACATGGC	TGCCCTGTAT	GTGGGCTCTG	ACCTGGAGAT
		TCATGTACCG			



1901	GGGGCCTGAC CCCCGGACTG				
1951	TGGATGGGCT ACCTACCCGA				
2001	GCTGCCTGAG CGACGGACTC			TGACATCCAG ACTGTAGGTC	
2051	GCAAGCTGAA CGTTCGACTT	CTGGGCCTCC GACCCGGAGG	CAAATCTACC GTTTAGATGG	CTGGCATCAA GACCGTAGTT	GGTGAGGCAG CCACTCCGTC
2101	CTGTGCAAGC GACACGTTCG	TGCTGAGGGG ACGACTCCCC	CACCAAGGCC GTGGTTCCGG	CTGACTGAGG GACTGACTCC	TGATCCCCCT ACTAGGGGGA
2151		GCTGAGCTGG CGACTCGACC	AGCTGGCTGA TCGACCGACT	GAACAGGGAG CTTGTCCCTC	ATCCTGAAGG TAGGACTTCC
2201				CCAAGGACCT GGTTCCTGGA	
2251	ATCCAGAAGC TAGGTCTTCG	AGGGCCAGGG TCCCGGTCCC	CCAGTGGACC GGTCACCTGG	TACCAAATCT ATGGTTTAGA	ACCAGGAGCC TGGTCCTCGG
2301				CAGGATGAGG GTCCTACTCC	
2351	CCAATGATGT GGTTACTACA	GAAGCAGCTG CTTCGTCGAC	ACTGAGGCTG TGACTCCGAC	TGCAGAAGAT ACGTCTTCTA	CACCACTGAG GTGGTGACTC
2401	TCCATTGTGA AGGTAACACT	TCTGGGGCAA AGACCCCGTT	GACCCCCAAG CTGGGGGTTC	TTCAAGCTGC AAGTTCGACG	CCATCCAGAA GGTAGGTCTT
2451					ACCTGGATCC TGGACCTAGG
2501					GTGGTACCAG CACCATGGTC
2551	CTGGAGAAGG GACCTCTTCC	AGCCCATTGT TCGGGTAACA	GGGGGCTGAG	ACCTTCTATO TGGAAGATAC	TGGCTGGGGC ACCGACCCCG
2601	TGCCAACAGG ACGGTTGTCC	GAGACCAAGC CTCTGGTTCG	TGGGCAAGGC	TGGCTATGTG	ACCAACAGGG TGGTTGTCCC
2651	GCAGGCAGAA CGTCCGTCTI	GGTGGTGACC	CTGACTGACA GACTGACTGA	CCACCAACCA CGTGGTTGGT	GAAGACTGCC CTTCTGACGG
2701	CTCCAGGCCA GAGGTCCGGT	TCTACCTGGC AGATGGACCG	CCTCCAGGAC GGAGGTCCTC	TCTGGCCTGG AGACCGGACG	AGGTGAACAT TCCACTTGTA
2751	TGTGACTGCC ACACTGACGC	TCCCAGTATO AGGGTCATAO	CCCTGGGCAT	CATCCAGGCO	C CAGCCTGATC G GTCGGACTAG



2851	GAGAAGGTGT		GGTGCCTGCC CCACGGACGG		
2901	TGAGCAGGTG	GACAAGCTGG	TGTCTGCTGG	CATCAGGAAG	GTGCTGTTCC
	ACTCGTCCAC	CTGTTCGACC	ACAGACGACC	GTAGTCCTTC	CACGACAAGG
2951	TGGATGGCAT				
	ACCTACCGTA	ACTGTTCCGG	GTCCTACTCG	TACTCTTCAT	GGTGAGGTTG
3001	TGGAGGGCTA	TGGCCTCTGA	CTTCAACCTG	CCCCCTGTGG	TGGCTAAGGA
	ACCTCCCGAT	ACCGGAGACT	GAAGTTGGAC	GGGGGACACC	ACCGATTCCT
3051	GATTGTGGCC	TCCTGTGACA	AGTGCCAGCT	GAAGGGGGAG	GCCATGCATG
3031			TCACGGTCGA		
3101			GGCATCTGGC		
			CCGTAGACCG		
3151			GGCTGTGCAT		
	CTCCCGTTCC	ACTAGGACCA	CCGACACGTA	CACCGGAGGC	CGATGTAACT
3201	GGCTGAGGTG	ATCCCTGCTG	AGACAGGCCA	GGAGACTGCC	TACTTCCTGC
•	CCGACTCCAC	TAGGGACGAC	TCTGTCCGGT	CCTCTGACGG	ATGAAGGACG
3251	TGAAGCTGGC	TGGCAGGTGG	CCTGTGAAGA	CCATCCACAC	TGCCAATGGC
			GGACACTTCT		ACGGTTACCG
3301	тосььсттсь	CTGGGGCCAC	AGTGAGGGCT	GCCTGCTGGT	GGGCTGGCAT
3301			TCACTCCCGA		
3351	CAAGCAGGAG	TTTGGCATCC	CCTACAACCC	CCAGTCCCAG	GGGGTGGTGG
JJJ1			GGATGTTGGG		
2401					
3401	·CCTCCATGAA				
			TTCTTCTAGT		
3451			TGTGCAGATG		
			ACACGTCTAC		
3501	CAAGAGGAAG				
	GTTCTCCTTC	CCCCCGTAGC	CCCCGATGAG	GCGACCCCTC	TCCTAACACC
3551	ACATCATTGC	CACAGACATC	CAGACCAAGG	AGCTCCAGAA	GCAGATCACC
					CGTCTAGTGG
3601	AAGATCCAGA	ACTTCAGGGT	GTACTACAGG	GACTCCAGGA	ACCCCCTGTG
3001					TGGGGGACAC
3651	GAAGGGCCCT	כרר אבריזיני	ጥርጥርር እ እርርር	CC A CC CC CC C T	GTGGTGATCC
2021					CACCACTAGG
2201	>00>0>	#C3C3#C3*C	. השרהשריריה	CCACCAACCC	CAAGATCATC
3701					GTTCTAGTAG
	TUUTGITGAG	ACTOTACTIC	CACCACGGGI	CCICCIICCG	GIICINGING



3801		GCAGATCTGC CGTCTAGACG	
3851	•	 CCCGTGCCTT GGGCACGGAA	
3901.		ATAAAATGAG TATTTTACTC	
3951		TGGGGGGTGG ACCCCCCACC	
4001	·	AGCAGGCATG TCGTCCGTAC	
4051		ACTGAAATGT TGACTTTACA	
4101		GGGTCTTATG CCCAGAATAC	
4151		GCACCAACTC CGTGGTTGAG	
4201		ATGCCCCCAT TACGGGGGTA	
4251		TGGTCGCCCC ACCAGCGGGG	
4301		TGTCTGGAAC ACAGACCTTG	
4351		GCAGCCACCG CGTCGGTGGC	TGTGACTGAC ACACTGACTG
4401			GTTCATCCGC CAAGTAGGCG
4451			TCTTTGACCC AGAAACTGGG
4501			CCAGCAGGTT GGTCGTCCAA
4551			ACATAAATAA TGTATTTATT
4601			TGCTGTCTTT ACGACAGAAA
4651			GTCTCGGTCG CAGAGCCAGC

Figure 26E

PCT/US01/28861 WO 02/022080

4751	GTTCAGATAC CAAGTCTATG				
4801	GCAGAGCTTC CGTCTCGAAG		GTGGTGTTGT CACCACAACA		
4851	GAGCGCTGGG CTCGCGACCC				
4901	CAGGGGCAGG GTCCCCGTCC				
4951	GGTGCATACG CCACGTATGC				
5001	GCTATGTTCC CGATACAAGG				
5051	CAGCACAGTG GTCGTGTCAC		ACTTGGGAAA TGAACCCTTT		
5101			GAGACGCCCT CTCTGCGGGA		
5151			GGCAATGGGC CCGTTACCCG		
5201					AGGATGAGAT TCCTACTCTA
5251					AGACTGCGGT TCTGACGCCA
5301					AGATTTGCAT TCTAAACGTA
5351					TGCGGGGCGA ACGCCCCGCT
5401	TGAAGAAAAC ACTTCTTTTG	GGTTTCCGGG CCAAAGGCCC	GTAGGGGAGA CATCCCCTCT	TCAGCTGGGA AGTCGACCCT	AGAAAGCAGG TCTTTCGTCC
5451	TTCCTGAGCA AAGGACTCGT				AAATCACACC TTTAGTGTGG
5501	TATTACCGGC ATAATGGCCG				CCGTCATCCC GGCAGTAGGG
5551	TGAGCAGGGG ACTCGTCCCC	GGCCACTTCG CCGGTGAAGC	TTAAGCATGT AATTCGTACA	CCCTGACTCG	CATGTTTTCC GTACAAAAGG
5601	CTGACCAAAT GACTGGTTTA				GCAGTTCTTG CGTCAAGAAC

Figure 26 F

5701				GGTCCCACAG CCAGGGTGTC	
5751				CCTCGTTTCG GGAGCAAAGC	
5801				TCGTCCAGAC AGCAGGTCTG	
5851				CAGCGTAGTC GTCGCATCAG	
5901				CCAGGGTGCG GGTCCCACGC	
5951				TCGCCCTGCG AGCGGGACGC	
6001				CCCCTCCGCG GGGGAGGCGC	
6051	TGGCGCGCAG ACCGCGCGTC	CTTGCCCTTG GAACGGGAAC	GAGGAGGCGC CTCCTCCGCG	CGCACGAGGG GCGTGCTCCC	GCAGTGCAGA CGTCACGTCT
6101	CTTTTGAGGG GAAAACTCCC	CGTAGAGCTT GCATCTCGAA	GGGCGCGAGA CCCGCGCTCT	AATACCGATT TTATGGCTAA	CCGGGGAGTA GGCCCCTCAT
6151	GGCATCCGCG CCGTAGGCGC	CCGCAGGCCC GGCGTCCGGG	CGCAGACGGT GCGTCTGCCA	CTCGCATTCC GAGCGTAAGG	ACGAGCCAGG TGCTCGGTCC
6201	TGAGCTCTGG ACTCGAGACC	CCGTTCGGGG GGCAAGCCCC	TCAAAAACCA AGTTTTTGGT	GGTTTCCCCC	ATGCTTTTTG TACGAAAAAC
6251	ATGCGTTTCT TACGCAAAGA	TACCTCTGGT	TTCCATGAGO AAGGTACTCG	CGGTGTCCAC GCCACAGGTG	GCTCGGTGAC CGAGCCACTG
6301					CTGTCCTCGA GACAGGAGCT
6351	GCGGTGTTCC CGCCACAAGG	GCGGTCCTCC	TCGTATAGAA AGCATATCTT	ACTCGGACCA TGAGCCTGGT	CTCTGAGACA GAGACTCTGT
6401	AAGGCTCGCG TTCCGAGCGC	TCCAGGCCAG AGGTCCGGTC	CACGAAGGAG GTGCTTCCTC	GCTAAGTGGG CGATTCACCC	AGGGGTAGCG TCCCCATCGC
6451	GTCGTTGTCC CAGCAACAGG	ACTAGGGGGT TGATCCCCCA	CCACTCGCTC	CAGGGTGTGA GTCCCACACT	AGACACATGT TCTGTGTACA
6501	CGCCCTCTTC GCGGGAGAA	GGCATCAAGG GCCGTAGTTCG	AAGGTGATTO	GTTTGTAGGT CAAACATCCA	GTAGGCCACG CATCCGGTGC
6551	TGACCGGGTG ACTGGCCCAC	TTCCTGAAGO	GGGGCTATA	AAGGGGGTGG	GGGCGCGTTC CCCGCGCAAG

Figure 266

6651	AGTACTCCCT	CTGAAAAGCG	GGCATGACTT	CTGCGCTAAG	ATTGTCAGTT
	TCATGAGGGA	GACTTTTCGC	CCGTACTGAA	GACGCGATTC	TAACAGTCAA
6701	TCCAAAAACG	ACCACCATTT	GATATTCACC	TGGCCCGCGG	TGATGCCTTT
6701			CTATAAGTGG		
6751			GGTCAGAAAA		
	CTCCCACCGG	CGTAGGTAGA	CCAGTCTTTT	CTGTTAGAAA	AACAACAGTT
6801	GCTTGGTGGC	AAACGACCCG	TAGAGGGCGT	TGGACAGCAA	CTTGGCGATG
	CGAACCACCG	TTTGCTGGGC	ATCTCCCGCA	ACCTGTCGTT	GAACCGCTAC
6851	GAGCGCAGGG	TTTGGTTTTT	GTCGCGATCG	GCGCGCTCCT	TGGCCGCGAT
0031			CAGCGCTAGC		
6901			GCGCAACGCA		
			CGCGTTGCGT		
6951	TGGTGCGCTC	GTCGGGCACC	AGGTGCACGC	GCCAACCGCG	GTTGTGCAGG
			TCCACGTGCG		
7001	GTGACAAGGT	CAACGCTGGT	GGCTACCTCT	CCGCGTAGGC	GCTCGTTGGT
	CACTGTTCCA	GTTGCGACCA	CCGATGGAGA	GGCGCATCCG	CGAGCAACCA
Ż051	CCAGCAGAGG	CGGCCGCCCT	TGCGCGAGCA	GAATGGCGGT	AGGGGGTCTA
, , , ,			ACGCGCTCGT		
7101	GCTGCGTCTC	GTCCGGGGGG	TCTGCGTCCA	CGGTAAAGAC	CCCGGGCAGC
,101			AGACGCAGGT		
7151	AGGCGCGCGT	CGAAGTAGTC	TATCTTGCAT	CCTTGCAAGT	CTAGCGCCTG
1231					GATCGCGGAC
7201					AGTGGGGGAC
					TCACCCCCTG
7251					GCAAATGTCG
					CGTTTACAGC
7301	TAAACGTAGA	GGGGCTCTCI	GAGTATTCCA	AGATATGTAG	GGTAGCATCT
	ATTTGCATCT	CCCCGAGAGA	CTCATAAGGT	TCTATACATO	CCATCGTAGA
7351	TCCACCGCGG	ATGCTGGCGC	GCACGTAATC	GTATAGTTC	TGCGAGGGAG
	AGGTGGCGCC	TACGACCGCG	CGTGCATTAC	CATATCAAGO	ACGCTCCCTC
7401	CGAGGAGGTC	GGGACCGAGG	TTGCTACGG	CGGGCTGCTC	TGCTCGGAAG
	GCTCCTCCAG	CCCTGGCTCC	AACGATGCCC	C GCCCGACGAC	ACGAGCCTTC
7451	ልሮጥልጥሮጥርርር	TGAAGATGG	ATGTGAGTT	GATGATATG	TTGGACGCTG
,451	TGATAGACGG	ACTTCTACCO	TACACTCAA	CTACTATACO	AACCTGCGAC
7501	ርልልርልቦርጥጥር	AAGCTGGCG	CTGTGAGAC	TACCGCGTC	A CGCACGAAGG
,501	CTTCTGCAAC	TTCGACCGC	A GACACTCTG	ATGGCGCAG	r GCGTGCTTCC

Figure 26 H

7601		AGTAGTCCAG TCATCAGGTC			
7651		TTCCACAGCT AAGGTGTCGA			
7701		TTGGATCGGA AACCTAGCCT			
7751		ACTGGTTGAC TGACCAACTG			
7801		TATGCCTGCG ATACGGACGC			
7851		CCTGACCATG GGACTGGTAC			
7901		CGCCCTGCTC GCGGGACGAG			
7951		GGCAGGGCGA CCGTCCCGCT			
8001		AAAGTTGCGT TTTCAACGCA			
8051		TTACCTGGGC AATGGACCCG			
8101		ACAATGTAAA TGTTACATTT			
8151		TTTAAGTTCC AAATTCAAGG			
8201		AAAGGGCCCA TTTCCCGGGT			AAGCGACGAA TTCGCTGCTT
8251	TGAGCTCCAC ACTCGAGGTG	AGGTCACGGG TCCAGTGCCC	CCATTAGCAT GGTAATCGTA	TTGCAGGTGG AACGTCCACC	TCGCGAAAGG AGCGCTTTCC
8301					GCAGTAGAAG CGTCATCTTC
8351					CGGCTAGGTC GCCGATCCAG
8401					ATGACCAGCA TACTGGTCGT
8451					ATAGGTCTCT TATCCAGAGA

Figure 26I

8551	GAAGAACTGG	ATCTCCCGCC	ACCAATTGGA	GGAGTGGCTA	TTGATGTGGT
•				CCTCACCGAT	
	C1.0110				
0.601	CARACERCAR	רתיייייייייייייייייייייייייייייייייייי	CCCCCCAAC	ACTCGTGCTG	CCTTTTCTAA
8601					
	CTTTCATCTT	CAGGGACGCT	GCCCGGCTTG	TGAGCACGAC	CGAAAACATT
8651				GGCTGTACAT	
	TTTGCACGCG	TCATGACCGT	CGCCACGTGC	CCGACATGTA	GGACGTGCTC
					•
8701	CTTCACCTGA	CGACCGCGCA	CAAGGAAGCA	GAGTGGGAAT	TTGAGCCCCT
0,01				CTCACCCTTA	
	CAACIGGACI	001000001	0		
		0mmm000m00	mccmcmmcm*	CTTCGGCTGC	TO THE PARTY OF TH
8751					
	GCGGACCGCC	CAAACCGACC	ACCAGAAGAT	GAAGCCGACG	AACAGGAACT
				•	
8801				GATCGGACCA	
	GGCAGACCGA	CGAGCTCCCC	TCAATGCCAC	CTAGCCTGGT	GGTGCGGCGC
8851	CGAGCCCAAA	GTCCAGATGT	CCGCGCGCGG	CGGTCGGAGC	TTGATGACAA
0031				GCCAGCCTCG	
	6010666111	chooleinen		000000	
		AMOCO A COMO	ጥ ር ር እ ጥ ር ር ጥር ጥ	GGAGCTCCCG	CCCCCTCACC
8901					
	GTAGCGCGTC	TACCCTCGAC	AGGTACCAGA	CCTCGAGGGC	GCCGCAGTCC
8951				CATAGACGGG	
	AGTCCGCCCT	CGAGGACGTC	CAAATGGAGC	GTATCTGCCC	AGTCCCGCGC
9001	GGCTAGATCC	AGGTGATACC	TAATTTCCAG	GGGCTGGTTG	GTGGCGGCGT
,				CCCGACCAAC	
	cconicino				
9051	CCARCCCTTC	CANCAGGGGG	CATCCCCCC	GCGCGACTAC	GGTACCGCGC
9051				CGCGCTGATG	
	GCTACCGAAC	GTTCTCCGGC	GIAGGGGGG	CGCGCIGNIG	CCAIGGCGCG
					00000
9101	GGCGGGCGGT				CTAAAAGCGG
	CCGCCCGCCA	CCCGGCGCCC	CCACAGGAAC	CTACTACGTA	GATTTTCGCC
					•
9151	TGACGCGGGC	GAGCCCCCGG	AGGTAGGGGG	GGCTCCGGAC	CCGCCGGGAG
					GGCGGCCCTC
				•	
0201	ACCCCCCACC	CCCACCTCCC	רפררפרפרפר	GGGCAGGAGC	TGGTGCTGCG
,	macacacacac	CCCTCCACCC	CCCCCCCCC	CCCCTCCTCG	ACCACGACGC
	TCCCCCGTCC	CCGIGCAGCC	GCGGCGCGCG	CCCG1CC1CG	Accidence
			6001 001 000		
9251	CGCGTAGGTT	GCTGGCGAAC	GCGACGACGC	GGCGGTTGAT	CTCCTGAATC
	GCGCATCCAA	CGACCGCTTG	CGCTGCTGCG	CCGCCAACTA	GAGGACTTAG
9301	TGGCGCCTCT	GCGTGAAGAC	GACGGGCCCG	GTGAGCTTGA	ACCTGAAAGA
	ACCGCGGAGA	CGCACTTCTG	CTGCCCGGGC	CACTCGAACT	TGGACTTTCT
	3.0000000000				
0251	CACTTCCACA	GAATCAATTT	CGGTGTCGTT	GACGGCGGCC	TGGCGCAAAA
3331	GWG1 1 CGUCU	CTTACTTAAA	GCCACAGCAA	CTGCCGCCGG	ACCGCGTTTT
	CICAMOCIGI	CILAGIIAAA	Jeenengen		
				**************************************	GGCCATGAAC
9401	TCTCCTGCAC	GICICCIGAG	11GICITGAT	WOOCGWICIC	CCCCTACTO
	AGAGGACGTG	CAGAGGACTC	AACAGAACTA	TUUGUTAGAG	CCGGTACTTG

Figure 26 J

9501			TGCGGGCCAT ACGCCCGGTA		
9551	GGCCTCCCTC CCGGAGGGAG	GTTCCAGACG CAAGGTCTGC	CGGCTGTAGA GCCGACATCT	CCACGCCCCC GGTGCGGGGG	TTCGGCATCG AAGCCGTAGC
9601			CGCGAGATTG GCGCTCTAAC		
9651			GCTGAAAGAG CGACTTTCTC		
9701			TACATAACCC ATGTATTGGG		
9751			AAGGCGCTCC TTCCGCGAGG		
9801			AGTTGCGCGC TCAACGCGCG		
9851			GCGACAGTGT CGCTGTCACA		
9901			TTCTTCAATC AAGAAGTTAG		
9951			GCGGTGGGGG CGCCACCCCC		
10001					CTCCCCGCGG GAGGGGCGCC
10051	CGACGGCGCA GCTGCCGCGT	TGGTCTCGGT ACCAGAGCCA	GACGGCGCGG CTGCCGCGCC	CCGTTCTCGC	GGGGGCGCAG
10101					GGGGGGCTGC
10151	CATGCGGCAG GTACGCCGTC	GGATACGGCG CCTATGCCGC	CTAACGATGC GATTGCTACG	ATCTCAACAA TAGAGTTGTT	TTGTTGTGTA AACAACACAT
10201	GGTACTCCGC CCATGAGGCG	CGCCGAGGGA	CCTGAGCGAG	TCCGCATCGA AGGCGTAGCT	CCGGATCGGA GGCCTAGCCT
10251	AAACCTCTCG TTTGGAGAGC	AGAAAGGCGT TCTTTCCGCA	CTAACCAGTC GATTGGTCAG	ACAGTCGCAA TGTCAGCGTT	GGTAGGCTGA CCATCCGACT
10301	GCACCGTGGC CGTGGCACCG	GGGCGGCAGC GCCGCCGTCG	: 020200000 : 020200000	CGGGGTTGTT	TCTGGCGGAG AGACCGCCTC
10351	GTGCTGCTGA CACGACGACT	TGATGTAATT	AAAGTAGGCO	GTCTTGAGAC CAGAACTCTC	GGCGGATGGT GCCGCCTACCA

Figure 26 K

10451	CGGCCATGCC GCCGGTACGG		GGCGCAGGTC CCGCGTCCAG	
10501			TCTTCTCCTT AGAAGAGGAA	
10551	• • • • • • • • • • • • • • • • • • • •		GGCGGAGTTT CCGCCTCAAA	
10601			CGAAGCCCCT GCTTCGGGGA	•
10651	AGCAGGGCTA TCGTCCCGAT		GCTAATATGG CGATTATACC	
	CTGCGTGAGG . GACGCACTCC	 		
10751			CCATAACGGA GGTATTGCCT	
10801			TACCTGAGAC ATGGACTCTG	
10851	-		CCGCACCAGG GGCGTGGTCC	
10901			AGAGGGGCCA TCTCCCCGGT	
10951			ATAAGGCGAT TATTCCGCTA	
11001			GGCGGTGGTG CCGCCACCAC	
11051			GCAGCGGCAA CGTCGCCGTT	
11101	ATGGTCGGGA TACCAGCCCT		GCGCAATCGT CGCGTTAGCA	
11151	GACCGTGCAA CTGGCACGTT		CACTCTTCCG GTGAGAAGGC	
11201	GATAAATTCG CTATTTAAGC			GAGCCCCGTA CTCGGGGCAT
11251	TCCGGCCGTC AGGCCGGCAG			GTCGAACCCA CAGCTTGGGT
11301	GGTGTGCGAC CCACACGCTG			TTCCTTCCAG AAGGAAGGTC

71 gure 26 L

11401	AAGCGGTTAG TTCGCCAATC		GAAAGCATTA CTTTCGTAAT		
11451			GTTGAGTCGC CAACTCAGCG		
11501			GAACGGGGGT CTTGCCCCCA		
11551			CGGAAACAGG GCCTTTGTCC		
11601			TGCGGCAGAT ACGCCGTCTA		
11651			CAGACATGCA GTCTGTACGT		
11701			ATCCGCGGTT TAGGCGCCAA		
11751			GGGCCCGGCA CCCGGGCCGT		
11801			GGAGCGCCCT CCTCGCGGGA		
11851			GCGTGAGGCG CGCACTCCGC		
11901			AGGAGCCCGA TCCTCGGGCT		
11951			CGGCATGGCC		
,12001					GTCCCGCGCG
	GCGTGTGCAC	CGCCGGCGGC	TGGACCATTG	GCGTATGCTC	CAGACGGTGA
	TGGTCCTCTA	ATTGAAAGTT	TTTTCGAAAT	TGTTGGTGCA	GCGTACGCTT CGCATGCGAA
	CACCGCGCGC	TCCTCCACCG	ATATCCTGAC	TACGTAGACA	GGGACTTTGT CCCTGAAACA
	TTCGCGCGAC	CTCGTTTTGG	GTTTATCGTT	CGGCGAGTAC	G GCGCAGCTGT CGCGTCGACA
12251					G GGATGCGCTG CCTACGCGAC

7 igure 26 M

12351				CTTGAGCCTG GAACTCGGAC	
12401				TGGGCAAGTT ACCCGTTCAA	
12451				GACAAGGAGG CTGTTCCTCC	
12501				GCTTACCTTG CGAATGGAAC	
12551				AGGCCGTGAG TCCGGCACTC	
12601				CACAGCCTGC GTGTCGGACG	
12651				CGAGTCCTAC GCTCAGGATG	
12701				GCGCCCTGGA CGCGGGACCT	
12751				CGCGCTGGCA	
12801				CGAGCCAGAG GCTCGGTCTC	GACGGCGAGT CTGCCGCTCA
12851				GCAAGACGCA CGTTCTGCGT	ACGGACCCGG TGCCTGGGCC
12901					CTCCACGGAC
12951					CGCGCAATCC
13001	TGACGCGTTC ACTGCGCAAG	CGGCAGCAGC	CGCAGGCCAA GCGTCCGGTT	CCGGCTCTCC GGCCGAGAGG	GCAATTCTGG CGTTAAGACC
13051	AAGCGGTGGT TTCGCCACCA	ccceccecec	GCAAACCCCA CGTTTGGGGT	CGCACGAGAA CGCGTGCTCTT	CCACGACCGC
13101	ATCGTAAACG TAGCATTTGC	CGCTGGCCGA CCGACCGGCT	AAACAGGGCC	TAGGCCGGGC	ACGAGGCCGG TGCTCCGGCC
13151	CCTGGTCTAC GGACCAGATG	GACGCGCTGC CTGCGCGACG	TTCAGCGCGT	GGCTCGTTAC	AACAGCGGCA TTGTCGCCGT
13201					G CGAGGCCGTG GCTCCGGCAC

Figure 26 N

13301	 TTCCTGAGTA AAGGACTCAT			
13351	 CAACTTTGTG GTTGAAACAC			
13401	 AGGTGTACCA TCCACATGGT	-		
13451	CTGCAGACCG GACGTCTGGC			
13501	 GGGGGTGCGG CCCCCACGCC	• •	•	
13551	 CGCCCAACTC GCGGGTTGAG			
13601	 GGCAGCGTGT CCGTCGCACA			
13651	 CGAGGCCATA GCTCCGGTAT			
13701	CAAGTGTCAG GTTCACAGTC			
13751	ACCCTAAACT TGGGATTTGA			
13801	CAGTTTAAAC GTCAAATTTG			
13851	TGAGCCTTAA ACTCGGAATT			
13901	ATGACCGCGC TACTGGCGCG			TATGCCTCAA ATACGGAGTT
13951				CCCCCCCCC
14001				ACTGGCTACC TGACCGATGG
14051				GGTAACGATG CCATTGCTAC
14101				GCAACCGCAG CGTTGGCGTC
14151				CGCTGCGAAA GCGACGCTTT

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14251	CGCGGTCAGA	TGCTAGTAGC	CCATTTCCAA	GCTTGATAGG	GTCTCTTACC
	GCGCCAGTCT	ACGATCATCG	GGTAAAGGTT	CGAACTATCC	CAGAGAATGG
14301	AGCACTCGCA	CCACCGCCC	GCGCCTGCTG	GGCGAGGAGG	AGTACCTAAA
	TCGTGAGCGT	GGTGGGCGGG	CGCGGACGAC	CCGCTCCTCC	TCATGGATTT
14351	CAACTCGCTG	CTGCAGCCGC	AGCGCGAAAA	AAACCTGCCT	CCGGCATTTC
	GTTGAGCGAC	GACGTCGGCG	TCGCGCTTTT	TTTGGACGGA	GGCCGTAAAG
14401			CTAGTGGACA GATCACCTGT		
14451	TACGCGCAGG ATGCGCGTCC	AGCACAGGGA TCGTGTCCCT	CGTGCCAGGC GCACGGTCCG	CCGCGCCGC	CCACCCGTCG GGTGGGCAGC
14501	TCAAAGGCAC	GACCGTCAGC	GGGGTCTGGT	GTGGGAGGAC	GATGACTCGG
	AGTTTCCGTG	CTGGCAGTCG	CCCCAGACCA	CACCCTCCTG	CTACTGAGCC
14551	CAGACGACAG	CAGCGTCCTG	GATTTGGGAG	GGAGTGGCAA	CCCGTTTGCG
	GTCTGCTGTC	GTCGCAGGAC	CTAAACCCTC	CCTCACCGTT	GGGCAAACGC
14601	CACCTTCGCC	CCAGGCTGGG	GAGAATGTTT	AAAAAAAAA	AAAAGCATGA
	GTGGAAGCGG	GGTCCGACCC	CTCTTACAAA	TTTTTTTTA	TTTTCGTACT
14651	TGCAAAATAA	AAAACTCACC	AAGGCCATGG	CACCGAGCGT	TGGTTTTCTT
	ACGTTTTATT	TTTTGAGTGG	TTCCGGTACC	GTGGCTCGCA	ACCAAAAGAA
14701	GTATTCCCCT CATAAGGGGA	TAGTATGCGG ATCATACGCC	CGCGCGGCGA	TGTATGAGGA ACATACTCCT	AGGTCCTCCT TCCAGGAGGA
14751	CCCTCCTACG GGGAGGATGC	AGAGTGTGGT TCTCACACCA	GAGCGCGGCG CTCGCGCCGC	CCAGTGGCGG	CGGCGCTGGG
14801	TTCTCCCTTC	GATGCTCCCC	TGGACCCGCC	GTTTGTGCCI	CCGCGGTACC
	AAGAGGGAAG	CTACGAGGGG	ACCTGGGCGG	CAAACACGGA	GGCGCCATGG
14851	TGCGGCCTAC	CGGGGGGAGA	AACAGCATCC	GTTACTCTGA	GTTGGCACCC
	ACGCCGGATG	GCCCCCTC1	TTGTCGTAGG	CAATGAGACT	CAACCGTGGG
14901	CTATTCGACA	CCACCCGTGT	GTACCTGGTG	GACAACAAGT	CAACGGATGT
	GATAAGCTGT	CGTGGGCACA	CATGGACCAC	CTGTTGTTCA	GTTGCCTACA
14951	GGCATCCCTG CCGTAGGGAC	AACTACCAGA TTGATGGTCT	A ACGACCACAG T TGCTGGTGTC	GTTGAAAGA	ACCACGGTCA TGGTGCCAGT
15001	TTCAAAACA? AAGTTTTGT?	TGACTACAGO ACTGATGTCO	CCGGGGGAGG CCCCCCCCCCCCCCCCCCCCCCCCCCCC	CAAGCACACACACACACACACACACACACACACACACAC	A GACCATCAAT CTGGTAGTTA
15051	CTTGACGACO GAACTGCTGO	GGTCGCACT	G GGGCGGCGA(C CCCGCCGCTY	CTGAAAACC GACTTTTGG	A TCCTGCATAC I AGGACGTATG
15101	CAACATGCCA GTTGTACGG	A AATGTGAACO T TTACACTTG	G AGTTCATGT	TACCAATAA A ATGGTTATT	TTTAAGGCGC AAATTCCGCG

Figure 26 P

15151	GGGTGA JT				
			GGATGATTCC		
15201	TACGAGTGGG ATGCTCACCC				
15251			ACGCGATCGT TGCGCTAGCA		
15201	CCACACACAA	CCCCCTTCTC	GAAAGCGACA	TCCCCCTA A A	CTTTCACACC
15301			CTTTCGCTGT		
15351	CGCAACTTCA	GACTGGGGTT	TGACCCCGTC	ACTGGTCTTG	TCATGCCTGG
			ACTGGGGCAG		
15401			TCCATCCAGA		
	CCATATATGT	TTGCTTCGGA	AGGTAGGTCT	GTAGTAAAAC	GACGGTCCTA
15451			AGCCGCCTGA		
	CGCCCCACCT	GAAGTGGGTG	TCGGCGGACT	CGTTGAACAA	CCCGTAGGCG
15501	AAGCGGCAAC		GGGCTTTAGG CCCGAAATCC		
					•
15551	000100-1010		TGTTGGATGT ACAACCTACA		•
15.601			GGCGGGGGTG		
15601			CCGCCCCAC		
15651	AGTGGCAGCG	GCGCGGAAGA	GAACTCCAAC	GCGGCAGCCG	CGGCAATGCA
			CTTGAGGTTG		
15701			ATCATGCCAT		
	CGGCCACCTC	CTGTACTTGC	TAGTACGGTA	AGCGCCGCTG	TGGAAACGGT
15751			GCTGAGGCCG		
	GTGCCCGACT	CCTCTTCGCG	CGACTCCGGC	TTCGTCGCCG	GCTTCGACGG
15801	GCCCCCGCTG		GGTCGAGAAG CCAGCTCTTC		
15851	CAAACCCCTG GTTTGGGGAC		GCAAGAAACG CGTTCTTTGC		
15001	ATGACAGCAC		•		•
13901			ATGGCGTCGA		
15951	GGCGACCCTC	AGACCGGAAT	CCGCTCATGG	ACCCTGCTTT	GCACTCCTGA
					CGTGAGGACT
16001	CGTAACCTGC				
	GCATTGGACG	CCGAGCCTCG	TCCAGATGAC	CAGCAACGGT	CTGTACTACG
16051	AAGACCCCGT				
	TTCTGGGGCA	CTGGAAGGCG	AGGTGCGCGG	TCTAGTCGTT	GAAAGGCCAC

Figure 26 Q

16151	GGCCGTCTAC CCGGCAGATG		TCCGCCAGTT AGGCGGTCAA		
16201	TCAATCGCTT AGTTAGCGAA		CAGATTTTGG GTCTAAAACC		
16251	ATCACCACCG TAGTGGTGGC		CGTTCCTGCT GCAAGGACGA		
16301	ACCGCTGCGC TGGCGACGCG		GAGGAGTCCA CTCCTCAGGT		
16351			TACGTTTACA ATGCAAATGT		
16401			CACTTTTTGA GTGAAAAACT		
16451			GCTGGGGCCT CGACCCCGGA		
16501	AACCGCCCCG	GTTCTTCGCG	TCCGACCAAC AGGCTGGTTG	TGGGTCACGC	GCACGCGCCC
16551	GTGATGGCGC	GCGGGACCCC		GCGCCGCGT	GACCCGCGTG
16601	GTGGCAGCTA	CTGCGGTAGC	TGCGCCACCA	CCTCCTCCGC	CGCAACTACA GCGTTGATGT
16651	GCGGGTGCGG	CGGTGGTCAC	AGGTGTCACC	TGCGCCGGTA	TCAGACCGTG AGTCTGGCAC
16701	CACGCGCCTC	GGGCCGCGAT	ACGATTTTAC	TTCTCTGCCG	GGAGGCGCGT CCTCCGCGCA
16751	TCGTGCAGCG	GTGGCGGCGG	CTGGGCCGTG	ACGGCGGGTT	GCGCGCGCC
	GCCGGGACGA	ATTGGCGCGT	GCAGCGTGGC	CGGCTGCCCG	GGCCATGCGG
	CGGCGAGCTT	CCGACCGGCG	CCCATAACAG	TGACACGGGG	CCAGGTCCAG GGTCCAGGTC
	CGCTGCTCGC	CGGCGGCGTC	CTCGGCGCCG	GTAATCACGA	ATGACTCAGG TACTGAGTCC
	CAGCGTCCCC	GTTGCACATA	ACCCACGCGC	TGAGCCAATC	GCCGGACGCG
17001	GTGCCCGTGC CACGGGCACG	GCACCCGCCC GCGCGGGGGGGGGGGGGGGGGGGGGGGG	CCCGCGCAAC GGGCGCGTTG	TAGATTGCAA ATCTAACGTT	GAAAAAACTA CTTTTTTGAT

7igure 26 R

17101	CTATGTCCAA GATACAGGTT	GCGCAAAATC CGCGTTTTAG			
17151	GAGATCTATG CTCTAGATAC	GCCCCCGAA CGGGGGGCTT			
17201		GTCAAAAAGA CAGTTTTTCT			
17251	ACGAGGTGGA TGCTCCACCT	ACTGCTGCAC TGACGACGTG			
17301	AAAGGTCGAC TTTCCAGCTG	GCGTAAAACG CGCATTTTGC			
17351		GAGCGCTCCA CTCGCGAGGT			
17401		CGAGGACCTG GCTCCTGGAC			
17451		GAAAGCGGCA CTTTCGCCGT			
17501		ACACCTAGCC TGTGGATCGG			
17551	GGCGCGAACG	ACCGTCCGAA TGGCAGGCTT	CTTTTCGCGC	CGGATTTCGC	GCTCAGACCA
17601		CCACCGTGCA GGTGGCACGT			
17651	TCTACAGAAC		GGCACCTTGG	ACCCGACCTC	GGGCTCCAGG
17701	CGCACGCCGG	TTAGTTCGTC	CACCGCGGCC	CTGACCCGCA	GCAGACCGTG CGTCTGGCAC
	CTGCAAGTCT	ATGGGTGATG	GTCATCGTGG	TCATAACGGT	CCGCCACAGA GGCGGTGTCT
	CCCGTACCTC	TGTGTTTGCA	GGGGCCAACG	GAGTCGCCAC	GCGGATGCCG CGCCTACGGC
	GCCACGTCC	CCAGCGACGC	CGGCGCAGG	TCTGGAGATG	GGAGGTGCAA CCTCCACGTT
17901	ACGGACCCGT TGCCTGGGCA	GGATGTTTCG CCTACAAAGC	GCAAAGTCGC	9 GGGGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	CGCGCCGTTC CGCGCGCAAG
17951					GCCCTACATC CGGGATGTAG

Figure 265

18051		CTACCCGACG GATGGGCTGC			
18101		CAGCCCGTGC GTCGGGCACG			
18151		CAGGACCCTG GTCCTGGGAC			
18201		AGCCGGTCTT TCGGCCAGAA			
18251		TTCCCGGTGC AAGGGCCACG			
18301		CGGCCACGGC GCCGGTGCCG			
18351		GCGCGTCGCA CGCGCAGCGT			
18401	CCTTATTCCA GGAATAAGGT	CTGATCGCCG GACTAGCGGC	CGGCGATTGG GCCGCTAACC	CGCCGTGCCC	GGAATTGCAT CCTTAACGTA
18451		GCAGGCGCAG CGTCCGCGTC			
18501	TTTTTAGTTT	ATAAAAAGTC TATTTTTCAG	ACCTGAGAGT	GCGAGCGAAC	CAGGACATTG
18551	ATAAAACATC		TAGTTGAAAC	GCAGAGACCG	GGGCGCTGTG
18601	CCGAGCGCGG	GCAAGTACCC	TTTGACCGTT	CTATAGCCGT	CCAGCAATAT GGTCGTTATA
18651	GAGCGGTGGC CTCGCCACCG	GCCTTCAGCT CGGAAGTCGA	GGGGCTCGCT	GTGGAGCGGC CACCTCGCCG	TTAAAATTA AATTTTTAAT
	AGCCAAGGTG	GCAATTCTTG	ATACCGTCGT	TCCGGACCTT	CAGCAGCACA GTCGTCGTGT
	CCGGTCTACG	ACTCCCTATT	CAACTTTCTC	GTTTTAAAGG	AACAAAAGGT TTGTTTTCCA
	CCATCTACCG	GACCGGAGAC	CGTAATCGCC	CCACCACCTG	CTGGCCAACC GACCGGTTGG
	TCCGTCACGT	TTTATTCTAA	TTGTCATTCG	AACTAGGGGC	CCCTCCCGTA CGGAGGGCAT
18901	GAGGAGCCTC CTCCTCGGAG	CACCGGCCGT GTGGCCGGCA	GGAGACAGTG CCTCTGTCAC	TCTCCAGAGG AGAGGTCTCC	GGCGTGGCGA CCGCACCGCT

Figure 26T

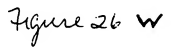
19001	GTACGAGGAG CATGCTCCTC		
19051	CCATGGCTAC GGTACCGATG		
19101	CCTCCCCCG GGAGGGGGC		
19151	CGTTGTTGTA GCAACAACAT		
19201	GTCCGCGATC CAGGCGCTAG		
19251	AACAGCATCG TTGTCGTAGC		
19301	 CTGATAGCTA GACTATCGAT		
19351	CAGAGGAGCT GTCTCCTCGA		
19401	TTCGATGATG AAGCTACTAC		
19451	CGGAGTACCT GCCTCATGGA		
19501	TACTTCAGCC ATGAAGTCGG		
19551	CGACGTGACC GCTGCACTGG		
19601	 TGGACCGTGA ACCTGGCACT		AGGCGCGGTT TCCGCGCCAA
19651			TCCACGTACT AGGTGCATGA
19701			GCCCTACTCT CGGGATGAGA
19751			ATCCTTGCGA TAGGAACGCT
19801			GAAGAGGACG CTTCTCCTGC
19851			AAAAACTCAC TTTTTGAGTG

Figure 26 U

19951	TCAAATAGGT		AAACACCTAA TTTGTGGATT		
20001	AACCTGAACC	TCAAATAGGA	GAATCTCAGT	GGTACGAAAC	AGAAATTAAT
			CTTAGAGTCA		
20051			AAAAAAGACT TTTTTTCTGA		
20101			CAAATGAAAA GTTTACTTTT		
20151			CTAGAAAGTC GATCTTTCAG		
20201			AGGCAATGGT TCCGTTACCA		
20251			TAGATATAGA ATCTATATCT		
20301			GAAGGTAACT CTTCCATTGA		
20351			TAATTACATT ATTAATGTAA		
20401			GCACGGGTAA CGTGCCCATT		
20451			GTTGTAGATT CAACATCTAA		
20501	CTTTCATACC GAAAGTATGG		TGATTCCATT ACTAAGGTAA		
20551	TTCTATGTGG AAGATACACC	AATCAGGCTG TTAGTCCGAC	TTGACAGCTA AACTGTCGAT	TGATCCAGAT ACTAGGTCTA	GTTAGAATTA CAATCTTAAT
20601	TTGAAAATCA AACTTTTAGT	TGGAACTGAA ACCTTGACTT	GATGAACTTC CTACTTGAAG	CAAATTACTG GTTTAATGAC	CTTTCCACTG GAAAGGTGAC
20651	GGÄGGTGTGA CCTCCACACT	TTAATACAGA AATTATGTCI	GACTCTTACC CTGAGAATGG	AAGGTAAAAC TTCCATTTTG	CTAAAACAGG GATTTTGTCC
20701	TCAGGAAAAT AGTCCTTTTA	GGATGGGAAA	AAGATGCTAC TTCTACGATG	AGAATTTTCA TCTTAAAAGT	GATAAAAATG CTATTTTTAC
20751	AAATAAGAGT TTTATTCTCA	TGGAAATAAT ACCTTTATTA	TTTGCCATGG	AAATCAATCT TTTAGTTAGA	AAATGCCAAC TTTACGGTTG
20801	CTGTGGAGAA GACACCTCTT	ATTTCCTGTA TAAAGGACAT	A CTCCAACATA TATOTTODAO	GCGCTGTATT CGCGACATA	TGCCCGACAA ACGGGCTGTT

Tigure 26 V

20901		GTGGTGGCTC CACCACCGAG	
20951		GTCCCTTGAC CAGGGAACTG	
21001		CTGGCCTGCG GACCGGACGC	
21051		TTCCACATCC AAGGTGTAGG	
21101		CCTGCCGGGC GGACGGCCCG	
21151		TGGTTCTGCA ACCAAGACGT	
21201		ATTAAGTTTG TAATTCAAAC	
21251		CAACACCGCC GTTGTGGCGG	
21301		 AGTCCTTTAA TCAGGAAATT	
21351		GCCAACGCTA CGGTTGCGAT	
21401		TTTCCGCGGC AAAGGCGCCG	
21451		TGGGCTCGGG ACCCGAGCCC	
21501		CTAGATGGAA GATCTACCTT	
21551			GGCCTGGCAA CCGGACCGTT
21601	TGACCGCCTG ACTGGCGGAC		TCAGTTGACG AGTCAACTGC
21651	GGGAGGGTTA CCCTCCCAAT	CAGTGTAACA GTCACATTGT	
21701			TCTATATCCC AGATATAGGG
21751			TTCCAGCCCA AAGGTCGGGT



21851		ACCAACACAA TGGTTGTGTT			
21901		GAAGGACAGG CTTCCTGTCC			
21051	The Coch beac	CGCAGTTGAC	»CC» TT» CCC	ACAAAAACTT	ጥርጥጥጥርርርልጥ
21951		GCGTCAACTG			
22001	CGCACCCTTT	GGCGCATCCC	ATTCTCCAGT	AACTTTATGT	CCATGGGCGC
22002		CCGCGTAGGG			
22051		CTGGGCCAAA			
•	TGAGTGTCTG	GACCCGGTTT	TGGAAGAGAT	GCGGTTGAGG	CGGGTGCGCG
22101		TTTTGAGGTG			
		AAAACTCCAC			
22151		AAGTCTTTGA TTCAGAAACT			
	CAAAACAAAC	TTCAGAAACT	GCACCAGGCA	CACGIGGICG	GCG1GGCGCC
22201		ACCGTGTACC			
	GCAGTAGCTT	TGGCACATGG	ACGCGTGCGG	GAAGAGCCGG	CCGTTGCGGT
22251		AAGCAAGCAA			
	GTTGTATTTC	TTCGTTCGTT	GTAGTTGTTG	TCGACGGCGG	TACCCGAGGT
22301	GTGAGCAGGA	ACTGAAAGCC	ATTGTCAAAG	ATCTTGGTTG	TGGGCCATAT
	CACTCGTCCT	TGACTTTCGG	TAACAGTTTC	TAGAACCAAC	ACCCGGTATA
22351					CTCCACACAA
	AAAAACCCGT	GGATACTGTT	CGCGAAAGGT	CCGAAACAAA	GAGGTGTGTT
22401					GGGGGCGTAC
	CGAGCGGACG	CGGTATCAGT	TATGCCGGCC	AGCGCTCTGA	CCCCCGCATG
22451					CTACCTCTTT
	TGACCTACCG	GAAACGGACC	TTGGGCGTGA	GTTTTTGTAC	GATGGAGAAA
22501	GAGCCCTTTG	GCTTTTCTGA	CCAGCGACTC	AAGCAGGTTI	ACCAGTTTGA
	CTCGGGAAAC	CGAAAAGACT	GGTCGCTGAG	TTCGTCCAAA	TGGTCAAACT
22551	GTACGAGTCA	CTCCTGCGCC	GTAGCGCCAT	TGCTTCTTCC	CCCGACCGCT
	CATGCTCAGT	GAGGACGCGG	CATCGCGGTA	ACGAAGAAGG	GGGCTGGCGA
22601	GTATAACGCT	GGAAAAGTCC	ACCCAAAGCG	TACAGGGGCC	CAACTCGGCC
	CATATTGCGA	CCTTTTCAGG	TGGGTTTCGC	: ATGTCCCCGG	GTTGAGCCGG
22651	GCCTGTGGAC	TATTCTGCTG	CATGTTTCTC	CACGCCTTTC	CCAACTGGCC
	CGGACACCTG	ATAAGACGAC	GTACAAAGAG	GTGCGGAAAC	GGTTGACCGG
22701	CCAAACTCCC	ATGGATCACA	ACCCCACCAT	GAACCTTATT	ACCGGGGTAC
	GGTTTGAGGG	TACCTAGTGT	TGGGGTGGT	CTTGGAATA	TGGCCCCATG

Figure 26 X

22801	CAGGAACAGE GTCCTTGTCG	TCTACAGCTT AGATGTCGAA	CCTGGAGUGU GGACCTCGCG	CACTUGULU. GTGAGCGGGA	TGAAGGCGTC
22851	CCACAGTGCG GGTGTCACGC	CAGATTAGGA GTCTAATCCT	GCGCCACTTC CGCGGTGAAG	TTTTTGTCAC AAAAACAGTG	TTGAAAAACA AACTTTTTGT
22901	TGTAAAAATA ACATTTTTAT	ATGTACTAGA TACATGATCT	GACACTTTCA CTGTGAAAGT	ATAAAGGCAA TATTTCCGTT	ATGCTTTTAT TACGAAAATA
22951	TTGTACACTC AACATGTGAG	TCGGGTGATT AGCCCACTAA	ATTTACCCCC TAAATGGGGG	ACCCTTGCCG TGGGAACGGC	TCTGCGCCGT AGACGCGGCA
23001	TTAAAAATCA AATTTTTAGT	AAGGGGTTCT TTCCCCAAGA	GCCGCGCATC CGGCGCGTAG	GCTATGCGCC CGATACGCGG	ACTGGCAGGG TGACCGTCCC
23051	ACACGTTGCG TGTGCAACGC	ATACTGGTGT TATGACCACA	TTAGTGCTCC AATCACGAGG	ACTTAAACTC TGAATTTGAG	AGGCACAACC TCCGTGTTGG
23101	TAGGCGCCGT	CGAGCCACTT	GTTTTCACTC CAAAAGTGAG	GTGTCCGACG	CGTGGTAGTG
23151	CAACGCGTTT GTTGCGCAAA	AGCAGGTCGG TCGTCCAGCC	GCGCCGATAT CGCGGCTATA	CTTGAAGTCG GAACTTCAGC	CAGTTGGGGC GTCAACCCCG
23201	CTCCGCCCTG GAGGCGGGAC	CGCGCGCGAG	TTGCGATACA AACGCTATGT	CAGGGTTGCA GTCCCAACGT	GCACTGGAAC CGTGACCTTG
23251	ACTATCAGCG TGATAGTCGC	CCGGGTGGTG GGCCCACCAC	CACGCTGGCC	AGCACGCTCT TCGTGCGAGA	TGTCGGAGAT ACAGCCTCTA
23301	GTCTAGGCGC	AGGTCCAGGA	GGCGCAACGA	GTCCCGCTTG	GGAGTCAACT CCTCAGTTGA
23351	AACCATCGAC	GGAAGGGTT	TTCCCGCGCA	. CGGGTCCGAP	TGAGTTGCAC ACTCAACGTG
23401	AGCGTGGCAT	CACCGTAGT	TTCCACTGGC	ACGGGCCAGA	GGGCGTTAGG CCCGCAATCC
	TATGTCGCG	G ACGTATITI	GGAACTAGAC	GAATTTTCG	ACCTGAGCCT TGGACTCGGA
•	AACGCGGAA	G TCTCTTCTT	G TACGGCGTTC	TGAACGGCC	A AAACTGATTG TTTGACTAAC
	CGGCCTGTC	C GGCGCAGCA	C GTGCGTCGT(GAACGCAGC(TGTTGGAGAT C ACAACCTCTA
	GACGTGGTG	T AAAGCCGGG	G TGGCCAAGAI	A GTGCTAGAA	G GCCTTGCTAG C CGGAACGATC
23651	ACTGCTCCT TGACGAGGA	T CAGCGCGCG A GTCGCGCGC	C TGCCCGTTT G ACGGGCAAA	r cgctcgtca a gcgagcagt	C ATCCATTTCA G TAGGTAAAGT

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WO 02/022080					PCT/US01/28861
23701	ATCACGTGCT	CCTTATTTAT	CATAATGCTT	CCGTGTAGAC	ACTTAAGCTC
23.02				GGCACATCTG	
23751	GCCTTCGATC			CAACGCGCAG GTTGCGCGTC	
23801	CGTGATGCTT	GTAGGTCACC	TCTGCAAACG	ACTGCAGGTA	CGCCTGCAGG
	GCACTACGAA	CATCCAGTGG	AGACGTTTGC	TGACGTCCAT	GCGGACGTCC
. 23851	AATCGCCCCA	TCATCGTCAC	AAAGGTCTTG	TTGCTGGTGA	AGGTCAGCTG
	TTAGCGGGGT	AGTAGCAGTG	TTTCCAGAAC	AACGACCACT	TCCAGTCGAC
23901	CAACCCGCGG				
	GTTGGGCGCC	ACGAGGAGCA	AGTCGGTCCA	GAACGTATGC	CGGCGGTCTC
23951	CTTCCACTTG	GTCAGGCAGT	AGTTTGAAGT	TCGCCTTTAG	ATCGTTATCC
	GAAGGTGAAC	CAGTCCGTCA	TCAAACTTCA	AGCGGAAATC	TAGCAATAGG
24001	ACGTGGTACT	TGTCCATCAG	CGCGCGCGCA	GCCTCCATGC	CCTTCTCCCA
24001				CGGAGGTACG	
0.405.	CGCAGACACG	> #GCCCO> G> G	mc> cccccmm	C) mc) cocm)	
24051				GTAGTGGCAT	
	GCGTCTGTGC	INGCCGIGIG	NOTCOCCCAN	GIAGIGGCAI	TANNO I GARA
24101	CCGCTTCGCT	GGGCTCTTCC	TCTTCCTCTT	GCGTCCGCAT	ACCACGCGCC
	GGCGAAGCGA	CCCGAGAAGG	AGAAGGAGAA	CGCAGGCGTA	TGGTGCGCGG
24151	ACTGGGTCGT	CTTCATTCAG	CCGCCGCACT	GTGCGCTTAC	CTCCTTTGCC
2.555	=			CACGCGAATG	
24201	ATGCTTGATT	AGCACCGGTG	GGTTGCTGAA	ACCCACCATT	TGTAGCGCCA
	TACGAACTAA	TCGTGGCCAC	CCAACGACTT	TGGGTGGTAA	ACATCGCGGT
24251	as mammamam	mmcmmccmcc	CDCDCCACCA	### CC#C#CC	#C>#C+CCC
24251				AATGGAGACC	TGATGGCGGG
24301	CGCTCGGGCT				
	GCGAGCCCGA	ACCCTCTTCC	CGCGAAGAAA	AAGAAGAACC	CGCGTTACCG
24351	CAAATCCGCC	GCCGAGGTCG	ATGGCCGCGG	GCTGGGTGTG	CGCGGCACCA
	GTTTAGGCGG	CGGCTCCAGC	TACCGGCGCC	CGACCCACAC	GCGCCGTGGT
24401	GCGCGTCTTG	TGATGAGTCT	TCCTCGTCCT	CGGACTCGAT	ACGCCGCCTC
24.00					TGCGGCGAG
24451	ATCCGCTTTT	TTGGGGGCGC	CCGGGGAGGC	GGCGGCGACG	GGGACGGGGA
27771					CCCTGCCCCT
24501	CGACACGTCC				
	GCTGTGCAGG	AGGTACCAAC	CCCCTGCAGC	GCGGCGTGGC	GCAGGCGCGA
24551	CGGGGGTGGT	TTCGCGCTGC	TCCTCTTCCC	GACTGGCCAT	TTCCTTCTCC
3 _					AAGGAAGAGG
				0101-01-0	
24601	TATAGGCAGA				
	ATATCCGTCT	TTTTCTAGTA	CUTCAGTCAG	CTCTTCTTCC	TGTCGGATTG

Figure 262 80/144

24701	CTACCACCTT GATGGTGGAA				
24751	ATCGAGCAGG TAGCTCGTCC				
24801	ACCAACAGAG TGGTTGTCTC			CAACGCAGAG GTTGCGTCTC	
24851	AACAAGTCGG TTGTTCAGCC			GCGACTACCT CGCTGATGGA	
24901	GACGACGTGC CTGCTGCACG			CAGTGCGCCA GTCACGCGGT	
24951	••••			CGCCATAGCG GCGGTATCGC	
25001				GCGTACCCCC CGCATGGGGG	
25051				CTCAACTTCT GAGTTGAAGA	
25101				CATCTTTTTC GTAGAAAAAG	
25151				GCCGAGCGGA CGGCTCGCCT	
25201				ATCGCCTCGC TAGCGGAGCG	
25251				CGAGAAGCGC GCTCTTCGCG	GCGGCAAACG CGCCGTTTGC
25301				GTCACTCTGG CAGTGAGACC	AGTGTTGGTG TCACAACCAC
25351					GCAGCATCGA CGTCGTAGCT
	CCAGTGGGTG	AAACGGATGG	GCCGTGAATT	GGATGGGGG	AAGGTCATGA TTCCAGTACT
25,451	GCACAGTCAT CGTGTCAGTA	GAGTGAGCTG CTCACTCGAC	ATCGTGCGCC TAGCACGCGG	GTGCGCAGCC	CCTGGAGAGG GGACCTCTCC
25501	GATGCAAATT CTACGTTTAA	TGCAAGAACA ACGTTCTTGT	AACAGAGGAG TTGTCTCCTC	GGCCTACCCG	CAGTTGGCGA GTCAACCGCT
25551	CGAGCAGCTA GCTCGTCGAT	GCGCGCTGGC CGCGCGACCG	TTCAAACGCG AAGTTTGCGC	CGAGCCTGCC	GACTTGGAGG CTGAACCTCC

7 igure 26 AA

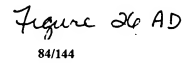
25651		GGTTCTTTGC CCAAGAAACG		
25701	AACATTGCAC TTGTAACGTG	TACACCTTTC ATGTGGAAAG		
25751		GGAGCTCTGC CCTCGAGACG		
25801		TTGGGCAAAA AACCCGTTTT		
25851	••	TACGTCCGCG ATGCAGGCGC		
25901		CATGGGCGTT GTACCCGCAA		
25951	AAGGAGCTGC TTCCTCGACG	AGAAACTGCT TCTTTGACGA		
26001		CGCTCCGTGG GCGAGGCACC	•	
26051		TAAAACCCTG ATTTTGGGAC		
26101		AGAACTTTAG TCTTGAAATC		
26151		TGCTGTGCAC ACGACACGTG		
26201	_	TCCGCCGCTT AGGCGGCGAA		
26251		CCTACCACTC GGATGGTGAG		
26301				CACCGCTCCC GTGGCGAGGG
26351				CGGTACCTTT GCCATGGAAA
26401				CGGGGTTGAA GCCCCAACTT
26451				TTTGTACCTG AAACATGGAC
26501				ATCCCGCCCG TAGGGCGGGC

Figure 26 AB

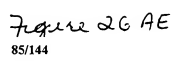
WO 02/022080					PCT/US01/28861
	CCTAA' 3G	AGCTTACCGC	CTGCGTCATT	ACCCA: C	
				TGGGTCCCGG	
26601	CCAATTGCAA	GCCATCAACA	AAGCCCGCCA	AGAGTTTCTG	CTACGAAAGG
	GGTTAACGTT	CGGTAGTTGT	TTCGGGCGGT	TCTCAAAGAC	GATGCTTTCC
26651	GACGGGGGGT				
				CGCTCCTCGA	
26701	CCCCCCCCC				
	GGGGGCGGCG	GCGTCGGGAT	AGTCGTCGTC	GGCGCCCGGG	AACGAAGGGT
26751	GGATGGCACC	CAAAAAGAAG	CTGCAGCTGC	CGCCGCCACC	CACGGACGAG
	CCTACCGTGG	GTTTTTCTTC	GACGTCGACG	GCGGCGGTGG	GTGCCTGCTC
26801	GAGGAATACT				
	CTCCTTATGA	CCCTGTCAGT	CCGTCTCCTC	CAAAACCTGC	TCCTCCTCCT
26851	GGACATGATG				
				GCTCCTTCGA	
26901	AAGAGGTGTC			GCCAGCGTAA	
26051	GCGCCCCAGA				
20931	••••			TACCGATGTT	
	000000101	1140000110	000100100		
27001	TCAGGCGCCG	CCGGCACTGC	CCGTTCGCCG	ACCCAACCGT	AGATGGGACA
	AGTCCGCGGC	GGCCGTGACG	GGCAAGCGGC	TGGGTTGGCA	TCTACCCTGT
27051	CCACTGGAAC				
				TCGGCGGCGG	
27101	GAGCAACAAC				
27151				ACCGCGCCCG	
2/151	CATAGTTGCT			GTTGTAGAGG	
27201	GCTTTCTTCT				
					GTAGGACGTA
· 27251	TACTACCGTC				
					CGCCGTCGTT
27301	CAGCAGCGGC				
					CTGAGACTGT
27351					GAGCGCTGCG
				-	CTCGCGACGC
27401					AACAGGATTT
	-				TTGTCCTAAA
27451					AGAACAAGAG TCTTGTTCTC
	WORD, I DWOW	CWINCOWINI	ANNOTTOICI	CG1CCCCG3	

7, guri 26. AC 83/144

27551	TCACAAAAGC AGTGTTTTCG			GCTGGAAGAC CGACCTTCTG	
27601				AGGACTAGTT TCCTGATCAA	
27651				CTCCAGCGGC GAGGTCGCCG	
27701	GCCAGCACCT CGGTCGTGGA			CAAGGAAATT GTTCCTTTAA	
27751				TTGCGGCTGG AACGCCGACC	
27801	••••			GCGGGACCCC	
27851				CCGAATTCTC GGCTTAAGAG	
27901				TTAATCCCCG AATTAGGGGC	
27951	CGACGGGACC	ACATGGTCCT	TTCAGGGCGA	CCCACCACTG GGGTGGTGAC	ACCATGAAGG
28001	GTCTCTGCGG	GTCCGGCTTC	AAGTCTACTG	TAACTCAGGG ATTGAGTCCC	CGCGTCGAAC
28051	GCCCGCCGAA	AGCAGTGTCC	CACGCCAGCG	CCGGGCAGGG	ATATTGAGTG
28101	GACTGTTAGT	CTCCCGCTCC	ATAAGTCGAG	TTGCTGCTCA	CGGTGAGCTC GCCACTCGAG
28151	GAGCGAACCA	GAGGCAGGCC	TGCCCTGTAA		CCGCGGCCGG
	CGAGAAGTAA	GTGCGGAGCA	GTCCGTTAGG	ATTGAGACGT	GACCTCGTCC CTGGAGCAGG
	AGACTCGGCG	CGAGACCTCC	GTAACCTTGA	GACGTTAAAT	TTGAGGAGTT AACTCCTCAA
	ACACGGTAGC	CAGATGAAAT	TGGGGAAGAG	CCCTGGAGGG	GGCCACTATC CCGGTGATAG
	GCCTAGTTAA	ATAAGGATTG	AAACTGCGCC	: ATTTCCTGAG	: GGCGGACGGC : CCGCCTGCCG
28401					TGAAACACCT ACTTTGTGGA



28451	GGTCCA、 JT CCAGGTGACA	CGCCGCCACA GCGGCGGTGT	AGTGCTTTGC TCACGAAACG	CCGCGA C GGCGCTGAGG	GGTGAGTTTT CCACTCAAAA
28501		ATTGCCCGAG TAACGGGCTC			
28551		CCCAGGGAGA GGGTCCCTCT			
28601	CCAGCGCCCC GGTCGCGGGG	CTGCTAGTTG GACGATCAAC	AGCGGGACAG TCGCCCTGTC	GGGACCCTGT CCCTGGGACA	GTTCTCACTG CAAGAGTGAC
28651		CTGTCCTAAC GACAGGATTG			
28701		AGTATAATAA TCATATTATT			
28751		TGTAAACGCC ACATTTGCGG			
28801		CTGGTACTTT GACCATGAAA			
28851		GACGGAGTGA CTGCCTCACT			
28901		AAAAAACACC TTTTTTGTGG			
28951	GCGTCACCGG CGCAGTGGCC	CCGCTGCACC GGCGACGTGG	ACACCTACCG TGTGGATGGC	CCTGACCGTA GGACTGGCAT	AACCAGACTT TTGGTCTGAA
29001	TTTCCGGACA AAAGGCCTGT	GACCTCAATA CTGGAGTTAT	ACTCTGTTTA TGAGACAAAT	CCAGAACAGG GGTCTTGTCC	AGGTGAGCTT
29051	AGAAAACCCT TCTTTTGGGA	TAGGGTATTA ATCCCATAAT	GGCCAAAGGC	GCAGCTACTG GCTCGATGAC	TGGGGTTTAT ACCCCAAATA
29101	GAACAATTCA CTTGTTAAGI	AGCAACTCTA TCGTTGAGAT	CGGGCTATTC	TAATTCAGGT ATTAAGTCCA	TTCTCTAGAA AAGAGATCTT
29151	TCGGGGTTGG AGCCCCAACG	GGTTATTCTC CCAATAAGAG	TGTCTTGTGA ACAGAACACI	TTCTCTTTAT AAGAGAAATA	TCTTATACTA AGAATATGAT
29201	ACGCTTCTCT TGCGAAGAGA	GCCTAAGGCT A CGGATTCCGA	CGCCGCCTGC	TGTGTGCACA ACACACGTGT	A TTTGCATTTA T AAACGTAAAT
29251	TTGTCAGCTT AACAGTCGA	TTTAAACGCT A AAATTTGCGA	GGGGTCGCCA CCCCAGCGGT	A CCCAAGATGA C GGGTTCTACT	TTAGGTACAT AATCCATGTA
29301	AATCCTAGG'	TTACTCACCO	TTGCGTCAGG AACGCAGTCG	CCACGGTACG GGTGCCATGG	ACCCAAAAGG TGGGTTTTCC
29351	TGGATTTTA ACCTAAAAT	A GGAGCCAGCO	TGTAATGTT	A CATTCGCAGO T GTAAGCGTCO	TGAAGCTAAT ACTTCGATTA



29451		AACAAAATTG TTGTTTTAAC		
29501		TACAGAGTAT ATGTCTCATA		
29551		TGTATACTTT ACATATGAAA		
29601	-	AAACAGTATA TTTGTCATAT		
29651		TTTCTGCTGC AAAGACGACG		
29701	-	TACTCTATAT ATGAGATATA		
29751		ATGCCTTAAT TACGGAATTA		
29801		CTCGCTGCTT GAGCGACGAA		
29851		GATTTAAACC CTAAATTTGG		
29901		TGACTCTATG ACTGAGATAC		
29951		CTGGATGTCA GACCTACAGT		
30001		CAGTCCAACT GTCAGGTTGA		
30051		CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC		ACAAATACAC TGTTTATGTG
30101				CATGTGGTGG GTACACCACC
30151		CGCTTATGTT GCGAATACAA		GGCTCATCTG CCGAGTAGAC
30201				CCCATCATTG GGGTAGTAAC
30251				ACTGAAACAC TGACTTTGTG
30301				TCCTCGAGTT AGGAGCTCAA

Figure 26 AF

30401		CACATCGAAG GTGTAGCTTC			
30451		ATTTGTCACC TAAACAGTGG			
30501		TTATCCAGTG AATAGGTCAC			
.30551		CATCCCCAGT GTAGGGGTCA			
30601		ATTATGAAAT TAATACTTTA			
30651		GTTTTGTTCC CAAAACAAGG			
30701		CTCGTATATG GAGCATATAC			
30751		GAAGCCTGGT CTTCGGACCA			
30801		CTTAGCCCTA GAATCGGGAT			
30851		ATGCCATGAA TACGGTACTT			
30901	AGGTGACGTT	CAAGTTGTTG GTTCAACAAC	GGCCGCCGAA	ACAGGGTCGG	TTAGTCGGAG
30951		TCCCACCCCC AGGGTGGGGG			TCTAACAGGA AGATTGTCCT
31001		GACACCCTAG CTGTGGGATC			ATTACAGAGC TAATGTCTCG
	TCGCGGACGA	TCTTTCTGCG	TCCCGTCGCC	GGCTCGTTGT	GCGCATGAAT CGCGTACTTA
	GTTCTCGAGG	TTCTGTACCA	ATTGAACGTG	GTCACGTTTI	GGGGTATCTT CCCCATAGAA
31151					ACCACCGGAC TGGTGGCCTG
31201					GGTGGTCATG CCACCAGTAC
31251					AAACCGAAGG TTTGGCTTCC

Figure 24 A6 87/144

31351		GATCTTATTC CTAGAATAAG		
31401		TAAAATCAGT ATTTTAGTCA		
31451		CCCTCCTCCC GGGAGGAGGG		
31501		CCACAATCTA GGTGTTAGAT		
31551		CCACTATCTT GGTGATAGAA		
31601		ACCTTCAACC TGGAAGTTGG		
31651		GCCTTTTCTT CGGAAAAGAA		
31701		CCCCTGGGGT GGGGACCCCA		
31751		GGCATGCTTG CCGTACGAAC		
31801	-	 CAACCTTACC GTTGGAATGG		
31851		CCAAGTCAAA GGTTCAGTTT		
31901		GAAGCCCTAA CTTCGGGATT		
31951		ACTCACCATG TGAGTGGTAC		
32001	CGTGCACGAC GCACGTGCTG	GCATTGCCAC CGTAACGGTG		
32051	CAGAAGGAAA GTCTTCCTTT	 CAAACATCAG GTTTGTAGTC	•	
32101	AGCAGTACCC TCGTCATGGG	TGCCTCACCC ACGGAGTGGG		
32151	TAGCTTGGGC ATCGAACCCG	AAGAGCCCAT TTCTCGGGTA		
32201	TAGGACTAAA ATCCTGATTT			CCTAAACACT GGATTTGTGA

Figure 26 AH

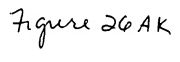
32301 .	AACTAAAGTT TTGATTTCAA	ACTGGAGCCT TGACCTCGGA	TGGGTTTTGA ACCCAAAACT	TTCACAAGGC AAGTGTTCCG	AATATGCAAC TTATACGTTG
32351			AGGATTGATT TCCTAACTAA		
32401			TGATGCTCAA ACTACGAGTT		
32451			TAAACTCAGC ATTTGAGTCG		
32501			TTTACAGCTT AAATGTCGAA		
32551			CAAGGGGTTG GTTCCCCAAC		
32601			GGCTTGAATT CCGAACTTAA		
32651			AAAATTGGCC TTTTAACCGG		ATTTGATTCA TAAACTAAGT
32701			ACTAGGAACT TGATCCTTGA		
32751					ACTTTGTGGA TGAAACACCT
32801					GAAAGATGCT CTTTCTACGA
32851					TTGCTACAGT AACGATGTCA
32901					GGAACAGTTC CCTTGTCAAG
32951	AAAGTGCTCA TTTCACGAGT	TCTTATTATA AGAATAATAT	AGATTTGACG TCTAAACTGC	AAAATGGAGI TTTTACCTCA	GCTACTAAAC CGATGATTTG
33001	AATTCCTTCC TTAAGGAAGG	TGGACCCAGA ACCTGGGTCT	ATATTGGAAC TATAACCTTG	TTTAGAAATG	GAGATCTTAC CTCTAGAATG
33051	TGAAGGCACA ACTTCCGTGT	GCCTATACAA CGGATATGTT	ACGCTGTTGG TGCGACAACC	ATTTATGCCT	AACCTATCAG TTGGATAGTC
33101	CTTATCCAAA GAATAGGTTT	A ATCTCACGGT TAGAGTGCCA	AAAACTGCCA A TTTTGACGGI	A AAAGTAACAT TTTCATTGTA	TGTCAGTCAA A ACAGTCAGTT
33151	GTTTACTTAJ CAAATGAAT	A ACGGAGACAI TGCCTCTGTT	A AACTAAACCT ADDTTTADTT 1	T GTAACACTAI A CATTGTGAT	A CCATTACACT CGTAATGTGA

Figure 26 AI

33251			GGCCACAACT CCGGTGTTGA		
33301	ACATCCTCTT	ACACTTTTTC	ATACATTGCC	CAAGAATAAA	GAATCGTTTG
	TGTAGGAGAA	TGTGAAAAAG	TATGTAACGG	GTTCTTATTT	CTTAGCAAAC
33351			ATTTTTCAAT		
			TAAAAAGTTA		
33401			CCCACCACCA GGGTGGTGGT		
33451	GTACCTTAAT	CAAACTCACA	GAACCCTAGT	ATTCAACCTG	CCACCTCCCT
			CTTGGGATCA		
33501			GTCCTTTCTC		
			CAGGAAAGAG		
33551			CATATTCTTA		
			GTATAAGAAT		
33601			CATCAGTGAT GTAGTCACTA		
			CTGTCCAGCT		
33651			GACAGGTCGA		
			GGGCGGCGAA		
33701	CCAACTTGCG	GTTGCTTAAC	CCCGCCGCTT	CCTCTTCAGG	TGCGGATGTA
				•	TGCTGCAGCA
33751					ACGACGTCGT
22001					GGAATACAAC
33801					CCTTATGTTG
22251		•			GCATAAGGCG
33851	TACCCTCACC	, ICICCICAGO	CTACTAACCG	ТОООООООТ	CGTATTCCGC
	INCCOTONCO	·	C11.C112.000		
33901	CCTTGTCCTC	CGGGCACAGC	AGCGCACCCT	GATCTCACTT	AAATCAGCAC
	GGAACAGGAG	GCCCGTGTCG	TCGCGTGGGA	CTAGAGTGAA	TTTAGTCGTG
33951	AGTAACTGC#	GCACAGCACC	ACAATATTGT	TCAAAATCCC	ACAGTGCAAG
					TGTCACGTTC
34001	GCGCTGTATC	CAAAGCTCAT	GGCGGGGACC	ACAGAACCCA	CGTGGCCATC
					GCACCGGTAG
34051	ATACCACAA	G CGCAGGTAGA	TTAAGTGGCG	ACCCCTCATA	AACACGCTGG
			•		TTGTGCGACC
34101	ACATAAACA?	TACCTCTTT	r GGCATGTTG1	AATTCACCAC	CTCCCGGTAC
	TGTATTTGT	A ATGGAGAAAJ	A CCGTACAACA	TTAAGTGGT	GAGGGCCATG

Figure 26 AJ

34201	GCTGGCCAAA	ACCTGCCCGC	CGGCTATACA	CTGCAGGGAA	CCGGGACTGG
	CGACCGGTTT	TGGACGGGCG	GCCGATATGT	GACGTCCCTT	GGCCCTGACC
34251	AACAATGACA	GTGGAGAGCC	CAGGACTCGT	AACCATGGAT	CATCATGCTC
74274				TTGGTACCTA	
	110111101	checicieo	0.0010		
34301	CTCATCATAT	CAATCTTCCC	ACAACACAGG	CACACGTGCA	TACACTTCCT
74707	CACTACTATA	CTTACAACC	TGTTGTGTCC	GTGTGCACGT	ATGTGAAGGA
	CAGIACIAIA	GIIACAACCO	70110.0100	0.0.0000	
34351	САССАТТАСА	ACCTCCTCCC	CCCTTAGAAC	CATATCCCAG	GGAACAACCC
24221				GTATAGGGTC	
	GICCIIEIIO.	200000000			
34401	ል ተጥርርጥርል ልጥ	САСССТАААТ	CCCACACTGC	AGGGAAGACC	TCGCACGTAA
34401				TCCCTTCTGG	
	IMOGRETIA	0.000			
34451	СТСАССТТСТ	GCATTGTCAA	AGTGTTACAT	TCGGGCAGCA	GCGGATGATC
34434				AGCCCGTCGT	
	GROT OCTUTE.				
34501	СТССАСТАТС	GTAGCGCGGG	TTTCTGTCTC	AAAAGGAGGT	AGACGATCCC
34304				TTTTCCTCCA	
	Quop : chine				
34551	тастстассс	AGTGCGCCGA	GACAACCGAG	ATCGTGTTGG	TCGTÄGTGTC
34332				TAGCACAACC	
34601	ATGCCAAATG	GAACGCCGGA	CGTAGTCATA	TTTCCTGAAG	CAAAACCAGG
	TACGGTTTAC				
34651	TGCGGGCGTG	ACAAACAGAT	CTGCGTCTCC	GGTCTCGCCG	CTTAGATCGC
• • • • • • • • • • • • • • • • • • • •				CCAGAGCGGC	
34701	TCTGTGTAGT	AGTTGTAGTA	TATCCACTCT	CTCAAAGCAT	CCAGGCGCCC
				GAGTTTCGTA	
34751	CCTGGCTTCG	GGTTCTATGT	AAACTCCTTC	ATGCGCCGCT	GCCCTGATAA
	GGACCGAAGC	CCAAGATACA	TTTGAGGAAG	TACGCGGCGA	CGGGACTATT
34801				GCCAACCTAC	
	GTAGGTGGTG	GCGTCTTATT	CGGTGTGGGT	CGGTTGGATG	TGTAAGCAAG
34851	TGCGAGTCAC	ACACGGGAGG	AGCGGGAAGA	GCTGGAAGAA	CCATGTTTTT
	ACGCTCAGTG	TGTGCCCTCC	TCGCCCTTCI	CGACCTTCTT	GGTACAAAAA
34901					ATCTATTAAG
	AAAAAATAAG	GTTTTCTAAT	AGGTTTTGGA	GTTTTACTTC	TAGATAATTC
34951	TGAACGCGCT	CCCCTCCGGT	GGCGTGGTC	AACTCTACAG	CCAAAGAACA
	ACTTGCGCGA	GGGGAGGCCA	CCGCACCAG	TTGAGATGTC	GGTTTCTTGT
35001	GATAATGGCA	TTTGTAAGAT	GTTGCACAA	GGCTTCCAA	AGGCAAACGG
	CTATTACCGT	AAACATTCTA	CAACGTGTT	CCGAAGGTTT	TCCGTTTGCC
		•		•	
35051					GTGAATCTCC
	GGGAGTGCAG	GTTCACCTGC	ATTTCCGAT	TGGGAAGTC	CACTTAGAGG



35151				CCGAATATTA GGCTTATAAT	
35201				CCTTCAGCCT GGAAGTCGGA	
35251				AGACCTGTAT TCTGGACATA	
35301				CGTAGGTCCC GCATCCAGGG	
35351				GACCAGCGCG CTGGTCGCGC	
35401	GCGGTCCTTG	GTACTGTTTT	CTTGGGTGTG	TGATTATGAC ACTAATACTG	TGCGTATGAG
35451				TAAGCTTGTT ATTCGAACAA	
35501	GCTATATTTT	ACGTTCCACG	ACGAGTTTTT	ATCAGGCAAA TAGTCCGTTT	CGGAGCGCGT
35551	TTTTTCTTTC	GTGTAGCATC	AGTACGAGTA	GCAGATAAAG CGTCTATTTC	CGTCCATTCG
35601	AGGCCTTGGT	GGTGTCTTTT	TCTGTGGTAA	TTTCTCTCAA AAAGAGAGTT	TGTACAGACG
35651	CCCAAAGACG	TATTTGTGTT	TTATTTTATT	CAAAAAAACA GTTTTTTTGT	AAATTTGTAA
35701	TCTTCGGACA	GAATGTTGTC	CTTTTTGTTG	GGAATATTCG	ATAAGACGGA TATTCTGCCT
35751	GATGCCGGTA	CGGCCGCACT	GGCATTTTTT	TGACCAGTGG	GTGATTAAAA CACTAATTTT
	TCGTGGTGGC	TGTCGAGGAG	CCAGTACAGG	CCTCAGTATT	TGTAAGACTC ACATTCTGAG
	CCATTTGTGT	AGTCCAACTA	AGTGTAGCCA	GTCACGATTT	AAGCGACCGA TTCGCTGGCT
	TTATCGGGCC	CCCTTATGTA	TGGGCGTCCG	CATCTCTGTT	CATTACAGCC
	GGGTATCCTC	CATATTGTTT	TAATTATCCI	CTCTTTTTGT	CATAAACACC GTATTTGTGG
36001	TGAAAAACCC ACTTTTTGGG	TCCTGCCTAG AGGACGGATC	GCAAAATAGC CGTTTTATCG	TGGGAGGGC	TCCAGAACAA AGGTCTTGTT

Figure 26 AL

36101	AAAGAAAACC TTTCTTTTGG			ACACGGCACC TGTGCCGTGG	
36151				GCGAGTATAT CGCTCATATA	
36201				AACACCCAGA TTGTGGGTCT	
36251				AACCCACAAC TTGGGTGTTG	
36301				TTCCCATTTT AAGGGTAAAA	
36351				CTAAAACCTA GATTTTGGAT	
36401				ACTCCACCCC TGAGGTGGGG	
					PacI
36451				ATTGATGATG TAACTACTAC	
36501	* ###	CCCACCCCAC	CCTCCATCCC	CTTCCCCATT	ATGATTCTTC
36201	TAAGCCTAGA	CGCTGCGCTC	CGACCTACCG	GAAGGGGTAA	TACTAAGAAG
36551	TCGCTTCCGG AGCGAAGGCC	CGGCATCGGG GCCGTAGCCC	ATGCCCGCGT TACGGGCGCA	TGCAGGCCAT ACGTCCGGTA	GCTGTCCAGG CGACAGGTCC
36601					AAAAGGCCAG TTTTCCGGTC
36651					CTCCGCCCCC
36701	CTGACGAGCA GACTGCTCGT	TCACAAAAAT AGTGTTTTTA	CGACGCTCAA GCTGCGAGTT	GTCAGAGGTG CAGTCTCCAC	GCGAAACCCG CGCTTTGGGC
36751					CCCTCGTGCG GGGAGCACGC
36801					GCCTTTCTCC GCGAAAGAGG
36851	CTTCGGGAAG GAAGCCCTTC	CGTGGCGCTT CGACCGCGAP	TCTCATAGCT AGAGTATCGA	CACGCTGTAC	GTATCTCAGT CATAGAGTCA
36901	TCGGTGTAGG AGCCACATCC	TCGTTCGCTC	CAAGCTGGGG GTTCGACCCG	TGTGTGCACGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	AACCCCCGT TTGGGGGGCA

Figure 26 AM

37001		CGACTTATCG GCTGAATAGC		
37051	***************************************	GGTATGTAGG CCATACATCC	 	
37101		TACACTAGAA ATGTGATCTT	 	
37151		CTTCGGAAAA GAAGCCTTTT	 	
37201		GTAGCGGTGG CATCGCCACC		
37251		GGATCTCAAG CCTAGAGTTC		
37301		GAACGAAAAC CTTGCTTTTG		
37351		TCTTCACCTA AGAAGTGGAT		
37401	-	GGTCTGACAG CCAGACTGTC		
37451		TGTCTATTTC ACAGATAAAG		
37501		TACGATACGG ATGCTATGCC		
37551		GAGACCCACG CTCTGGGTGC		
37601		GGAAGGGCCG CCTTCCCGGC		
37651		GTCTATTAAT CAGATAATTA		AAGTAGTTCG TTCATCAAGC
37701	CCAGTTAATA GGTCAATTAT			GCATCGTGGT CGTAGCACCA
37751				TCCCAACGAT AGGGTTGCTA
37801				GGTTAGCTCC CCAATCGAGG
37851	TTCGGTCCTC AAGCCAGGAG			TGTTATCACT ACAATAGTGA

Figure 26 AN

37951	GATGCTTTTC TGTGACTGGT GAGTACTCAA CCAAGTCATT CTGAGAATAG CTACGAAAAG ACACTGACCA CTCATGAGTT GGTTCAGTAA GACTCTTATC
38001	TGTATGCGGC GACCGAGTTG CTCTTGCCCG GCGTCAACAC GGGATAATAC ACATACGCCG CTGGCTCAAC GAGAACGGGC CGCAGTTGTG CCCTATTATG
38051	CGCGCCACAT AGCAGAACTT TAAAAGTGCT CATCATTGGA AAACGTTCTT GCGCGGTGTA TCGTCTTGAA ATTTTCACGA GTAGTAACCT TTTGCAAGAA
38101	CGGGGCGAAA ACTCTCAAGG ATCTTACCGC TGTTGAGATC CAGTTCGATG GCCCCGCTTT TGAGAGTTCC TAGAATGGCG ACAACTCTAG GTCAAGCTAC
38151	TAACCCACTC GTGCACCCAA CTGATCTTCA GCATCTTTTA CTTTCACCAG ATTGGGTGAG CACGTGGGTT GACTAGAAGT CGTAGAAAAT GAAAGTGGTC
38201	CGTTTCTGGG TGAGCAAAAA CAGGAAGGCA AAATGCCGCA AAAAAGGGAA GCAAAGACCC ACTCGTTTTT GTCCTTCCGT TTTACGGCGT TTTTTCCCTT
38251	TAAGGGCGAC ACGGAAATGT TGAATACTCA TACTCTTCCT TTTTCAATAT ATTCCCGCTG TGCCTTTACA ACTTATGAGT ATGAGAAGGA AAAAGTTATA
38301	TATTGAAGCA TTTATCAGGG TTATTGTCTC ATGAGCGGAT ACATATTTGA ATAACTTCGT AAATAGTCCC AATAACAGAG TACTCGCCTA TGTATAAACT
38351	ATGTATTTAG AAAAATAAAC AAATAGGGGT TCCGCGCACA TTTCCCCGAA TACATAAATC TTTTTATTTG TTTATCCCCA AGGCGCGTGT AAAGGGGCTT
38401	AAGTGCCACC TGACGTCTAA GAAACCATTA TTATCATGAC ATTAACCTAT TTCACGGTGG ACTGCAGATT CTTTGGTAAT AATAGTACTG TAATTGGATA
38451	AAAAATAGGC GTATCACGAG GCCCTTTCGT CTTCAAGAAT TGGATCCGAA TTTTTATCCG CATAGTGCTC CGGGAAAGCA GAAGTTCTTA ACCTAGGCTT
	PacI
38501	TTCTTAATTT CTTAATTAA (SEQ ID NO:32) AAGAATTAAA GAATTAATT (SEQ ID NO:33)

Figure 26 AD

1	CATCATCAAT	AATATACCTT	ATTTTGGATT	GAAGCCAATA	TGATAATGAG
	GTAGTAGTTA	TTATATGGAA	TAAAACCTAA	CTTCGGTTAT	ACTATTACTC
	•				
51				TGGGAACGGG	
	CCCCACCTCA	AACACTGCAC	CGCGCCCCGC	ACCCTTGCCC	CGCCCACTGC
101				GTGTGGCGGA	
	ATCATCACAC	CGCCTTCACA	CTACAACGTT	CACACCGCCT	TGTGTACATT
151	CCCACCCATC	TYCCE A A A GT	CACCTTTTTC	GTGTGCGCCG	GTGTACACAG
131				CACACGCGGC	
	COCTOCCIAC	Accolline.	0.00.11111		
201				GATGTTGTAG	
	CTTCACTGTT	AAAAGCGCGC	CAAAATCCGC	CTACAACATC	ATTTAAACCC
•					
251				GGGAAAACTG	
	GCATTGGCTC	ATTCTAAACC	GGTAAAAGCG	CCCTTTTGAC	TTATTCTCCT
				m> GOGCCM> 3	m > mmm/cm/cm >
301	AGTGAAATCT	GAATAATTTT	GIGITACICA	TAGCGCGTAA ATCGCGCATT	TATTIGICIA TATTIGICIA
	TCACTTTAGA	CTTATTAAAA	CACAATGAGT	ATCGCGCATT	ATAMACAGAT
351	GGGCCGCGG	GACTTTGACC	GTTTACGTGG	AGACTCGCCC	AGGTGTTTTT
331	CCCGCCGCCC	CTGAAACTGG	CAAATGCACC	TCTGAGCGGG	TCCACAAAAA
	CCCGGCGCCC	C. C	•		•
401				TTGGCGTTTT	
	GAGTCCACAA	AAGGCGCAAG	GCCCAGTTTC	AACCGCAAAA	TAATAATATC
451	GCGGCCGCGA	TCCATTGCAT	ACGTTGTATC	CATATCATAA	TATGTACATT
	CGCCGGCGCT	AGGTAACGTA	TGCAACATAG	GTATAGTATT	ATACATGTAA
F 0 *	m> m> mmcccm	C	እጥጥ እርርርርር እ	TGTTGACATT	GATTATTGAC
501				ACAACTGTAA	
	MINIMACCON	GIACAGGIIG			•
551	TAGTTATTAA	TAGTAATCAA	TTACGGGGTC	ATTAGTTCAT	AGCCCATATA
					TCGGGTATAT
601	TGGAGTTCCG	CGTTACATAA	CTTACGGTAA	ATGGCCCGCC	TGGCTGACCG
	ACCTCAAGGC	GCAATGTATT	GAATGCCATT	TACCGGGCGG	ACCGACTGGC
			G1 CCDC1 1 D1	> mc > ccm > mc	カカンとともできてか
651	CCCAACGACC	CCCGCCCATI	GACGICAAIA	MACACCIAIG	TTCCCATAGT AAGGGTATCA
	GGGTTGCTGG	GGGCGGGTAA	CIGCAGIIAI	INCIGCAINC	ANGGGIATER
701	N N C C C C N N T N	CCCACTTCC	ATTGACGTCA	ATGGGTGGAG	TATTTACGGT
701	TTCCCCTTAT	CCCTGAAAGG	TAACTGCAGT	TACCCACCTC	ATAAATGCCA
	11000011111				
751	AAACTGCCCA	CTTGGCAGTA	CATCAAGTGT	ATCATATGCC	AAGTACGCCC
	TTTGACGGGT	GAACCGTCAT	GTAGTTCACA	TAGTATACGG	TTCATGCGGG
801	CCTATTGACG	TCAATGACGG	TAAATGGCCC	GCCTGGCATI	ATGCCCAGTA
	GGATAACTGC	AGTTACTGCC	ATTTACCGG	G CGGACCGTAA	TACGGGTCAT

Figure 27A

901	TCGCTATTAC AGCGATAATG				
951	TAGCGGTTTG ATCGCCAAAC				
1001	TGGGAGTTTG ACCCTCAAAC				
1051	ACAACTCCGC TGTŢGAGGCG				
1101	GTCTATATAA CAGATATATT				
1151	CCATCCACGC GGTAGGTGCG				
1201	AGGCGCCGGC	CCTTGCCACG	TAACCTTGCG	CCTAAGGGGC	
1251	GAGATCTGCC CTCTAGACGG	ACCATGGCCG TGGTACCGGC	GCAAGTGGTC CGTTCACCAG	CAAGAGGTCC GTTCTCCAGG	GTGCCCGGCT CACGGGCCGA
1301	CCAGGTGGCA	СТСССТСТСС	TACTCCTCCC	CCGAGCCCGC GGCTCGGGCG	GCGGCTGTCC
1351	CACTCCTCCT	GGCTCGGGCG	GCGTCACCCG	GTGGGCGCCG CACCCGCGGC	ACAGGTCCCT
1401	GGACCTCTTC	GTGCCGCGGT	AGTGGAGGAG	CAACACCGCC GTTGTGGCGG	CGGTGGTTGC
1451	GGCTGACGCG	GACCGACCTC	CGGGTCCTCC	ACGAGGAGGT TGCTCCTCCA	CCCGAAGGGG
1501	CACTCCGGGG	TCCACGGGGA	CTCCGGGTAC	TGGATGTTCC	GCGCCGTGGA CGCGGCACCT
1551		AAGGACTTCC	TCTTCCCGCC	GGACCTCCCG	GACTAGGTGA
1601	GGGTCTTCTC	CGTCCTGTAG	GACCTGGACA	CCCACATGGT	CACCCAGGGC
1651					TCAGGTTCCC AGTCCAAGGG
1701	CCTGACCTTC GGACTGGAAG				CCCGAGAAGG GGGCTCTTCC
1751					CCCCATGTCC GGGGTACAGG

Figure 27B

1851	CTCCAAGCTG GAGGTTCGAC				
1901	ACAAGGACTG TGTTCCTGAC	CTAAAGCCCG GATTTCGGGC			
1951		TTTGCCCCTC AAACGGGGAG			
2001	CACTCCCACT GTGAGGGTGA	GTCCTTTCCT CAGGAAAGGA			
2051		TCATTCTATT AGTAAGATAA			
2101		GGGAAGACAA CCCTTCTGTT			
2151		CGCCCCCCC			
2201		ATATAAGGTG TATATTCCAC			
2251		CGCCGCCATG GCGGCGGTAC			
2301		TGACAACGCG ACTGTTGCGC			
2351		TCCAGCATTG AGGTCGTAAC			GCAAACTCTA CGTTTGAGAT
2401		CTACGAGACC GATGCTCTGG			
2451	TCCGCCGCCG AGGCGGCGGC	CTTCAGCCGC GAAGTCGGCG	TGCAGCCACC ACGTCGGTGG	GCCCGCGGA CGGGCGCCCT	TTGTGACTGA AACACTGACT
2501	CTTTGCTTTC GAAACGAAAG				CGTTCATCCG GCAAGTAGGC
2551	CCCGCGATGA GGGCGCTACT				TTCTTTGACC AAGAAACTGG
2601	CGGGAACTTA GCCCTTGAAT				GCCAGCAGGT CGGTCGTCCA
		TTCCGAAGGA	GGGGAGGGTT	ACGCCAAATT	TTGTATTTAT
2701					TTGCTGTCTT AACGACAGAA

Figure 27C

2751	TATTTAGGGGG	TTTTGCGCGC	GCGGTAGGCC	CGGGACCA	GGTCTCGGTC
2,31	ATAAATCCCC	AAAACGCGCG	CGCCATCCGG	GCCCTGGTCG	CCAGAGCCAG
2801	GTTGAGGGTC CAACTCCCAG	CTGTGTATTT GACACATAAA	TTTCCAGGAC AAAGGTCCTG	GTGGTAAAGG CACCATTTCC	TGACTCTGGA ACTGAGACCT
2851	TGTTCAGATA	CATGGGCATA	AGCCCGTCTC	TGGGGTGGAG	GTAGCACCAC
	ACAAGTCTAT	GTACCCGTAT	TCGGGCAGAG	ACCCCACCTC	CATCGTGGTG
2901	TGCAGAGCTT ACGTCTCGAA				
2951	GGAGCGCTGG	GCGTGGTGCC	TAAAAATGTC	TTTCAGTAGC	AAGCTGATTG
	CCTCGCGACC	CGCACCACGG	ATTTTTACAG	AAAGTCATCG	TTCGACTAAC
3001	CCAGGGGCAG	GCCCTTGGTG	TAAGTGTTTA ATTCACAAAT	CAAAGCGGTT	AAGCTGGGAT
3051			GAGATGCATC CTCTACGTAG		
	CCCACGTATG	CACCCCTATA	CICIACGIAG	AACCIGACAI	Manifella
3101	GGCTATGTTC	CCAGCCATAT	CCCTCCGGGG	ATTCATGTTG	TGCAGAACCA
	CCGATACAAG	GGTCGGTATA	GGGAGGCCCC	TAAGTACAAC	ACGTCTTGGT
3151	CCAGCACAGT	GTATCCGGTG	CACTTGGGAA	ATTTGTCATG	TAGCTTAGAA
	GGTCGTGTCA	CATAGGCCAC	GTGAACCCTT	TAAACAGTAC	ATCGAATCTT
3201	GGAAATGCGT	GGAAGAACTT	GGAGACGCCC	TTGTGACCTC	CAAGATTTTC
•••	CCTTTACGCA	CCTTCTTGAA	CCTCTGCGGG	AACACTGGAG	GTTCTAAAAG
3251	САТССАТТСС	TCCATAATGA	TGGCAATGGG	CCCACGGGCG	GCGGCCTGGG
3232	GTACGTAAGC	AGGTATTACT	ACCGTTACCC	GGGTGCCCGC	CGCCGGACCC
3301	CGAAGATATT	TCTGGGATCA	CTAACGTCAT	AGTTGTGTTC	CAGGATGAGA
	GCTTCTATAA	AGACCCTAGT	GATTGCAGTA	TCAACACAAG	GTCCTACTCT
3351	TCGTCATAGG	CCATTTTTAC	AAAGCGCGGG	CGGAGGGTGC	CAGACTGCGG
	AGCAGTATCC	GGTAAAAATG	TTTCGCGCCC	GCCTCCCACG	GTCTGACGCC
3401	TATAATGGTT	CCATCCGGCC	CAGGGGCGTA	GTTACCCTCA	CAGATTTGCA
	ATATTACCAA	GGTAGGCCGG	GTCCCCGCAT	CAATGGGAGI	GTCTAAACGT
3451	TTTCCCACGC	TTTGAGTTCA	GATGGGGGGA	TCATGTCTAC	CTGCGGGGCG
	AAAGGGTGCG	AAACTCAAGI	CTACCCCCCT	AGTACAGATO	GACGCCCCGC
3501	ATGAAGAAAA	CGGTTTCCGG	GGTAGGGGAG	ATCAGCTGG	AAGAAAGCAG
3301	TACTTCTTTT	GCCAAAGGCC	CCATCCCCTC	TAGTCGACCO	TTCTTTCGTC
3551	GTTCCTGAGC	AGCTGCGACT	TACCGCAGC	GGTGGGCCC	TAAATCACAC
	CAAGGACTCG	TCGACGCTG	A ATGGCGTCG(G CCACCCGGG(ATTTAGTGTG
3601	CTATTACCG	CTGCAACTG	TAGTTAAGA	AGCTGCAGC	GCCGTCATCC
	GATAATGGCC	GACGTTGAC	ATCAATTCT(TCGACGTCG	A CGGCAGTAGG
3651	CTGAGCAGG	GGGCCACTT	GTTAAGCAT	G TCCCTGACT	C GCATGTTTTC
	GACTCGTCC	CCCGGTGAA	G CAATTCGTAG	CAGGGACTGA	G CGTACAAAAG



3701	CCTGACCAAA GGACTGGTTT			GCCCAGCGAT CGGGTCGCTA	
3751	GCAAGGAAGC CGTTCCTTCG			GACCGTCCGC CTGGCAGGCG	
3801	CTTTTGAGCG GAAAACTCGC			CGGTCCCACA GCCAGGGTGT	
3851				TCCTCGTTTC AGGAGCAAAG	
3901		GACATGCCGT	CATCAGCCAC	GAGCAGGTCT	GCCCGGTCCC
3951	AGTACAGAAA	GGTGCCCGCG	TCCCAGGAGC	TCAGCGTAGT AGTCGCATCA	GACCCAGTGC
4001	CACTTCCCCA	CGCGAGGCCC	GACGCGCGAC	GCCAGGGTGC CGGTCCCACG	CGAACTCCGA
4051	CCAGGACGAC	CACGACTTCG	CGACGGCCAG	TTCGCCCTGC AAGCGGGACG	CGCAGCCGGT
4101	CCATCGTAAA	CTGGTACCAC	AGTATCAGGT	GCCCCTCCGC CGGGGAGGCG	CCGCACCGGG
4151	AACCGCGCGT	CGAACGGGAA	CCTCCTCCGC	CCGCACGAGG GGCGTGCTCC	CCGTCACGTC
4201	TGAAAACTCC	CGCATCTCGA	ACCCGCGCTC	AAATACCGAT TTTATGGCTA	AGGCCCCTCA
4251	TCCGTAGGCG	CGGCGTCCGG	GGCGTCTGCC	AGAGCGTAAG	
		CGGCAAGCCC	CAGTTTTTGG	TCCAAAGGGG	GTACGAAAAA
		AATGGAGACC	AAAGGTACTC	GGCCACAGGT	GCGAGCCACT
		CAGGCACAGG	GGCATATGTC	TGAACTCTCC	GGACAGGAGC
		GCGCCAGGAG	GAGCATATCT	TTGAGCCTGG	TGAGACTCTG
		CAGGTCCGGT	CGTGCTTCCT	CCGATTCACC	CTCCCCATCG
		GTGATCCCCC	AGGTGAGCGA	GGTCCCACAC	TTCTGTGTAC
4601	TCGCCCTCTT AGCGGGAGAA				TGTAGGCCAC ACATCCGGTG



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4701				CGAGGGCCAG GCTCCCGGTC	
4751				TCTGCGCTAA AGACGCGATT	
4801				CTGGCCCGCG GACCGGGCGC	
4851				AGACAATCTT TCTGTTAGAA	
4901				TTGGACAGCA AACCTGTCGT	
4951				GGCGCGCTCC CCGCGCGAGG	
5001	TGTTTAGCTG ACAAATCGAC	CACGTATTCG GTGCATAAGC	CGCGCAACGC GCGCGTTGCG	ACCGCCATTC TGGCGGTAAG	GGGAAAGACG CCCTTTCTGC
5051	GTGGTGCGCT CACCACGCGA	CGTCGGGCAC GCAGCCCGTG	CAGGTGCACG GTCCACGTGC	CGCCAACCGC GCGGTTGGCG	GGTTGTGCAG CCAACACGTC
5101	GGTGACAAGG CCACTGTTCC	TCAACGCTGG AGTTGCGACC	TGGCTACCTC ACCGATGGAG	TCCGCGTAGG AGGCGCATCC	CGCTCGTTGG GCGAGCAACC
5151	TCCAGCAGAG AGGTCGTCTC	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	TTGCGCGAGC AACGCGCTCG	AGAATGGCGG TCTTACCGCC	TAGGGGGTCT ATCCCCCAGA
5201	AGCTGCGTCT TCGACGCAGA	CGTCCGGGGG GCAGGCCCCC	GTCTGCGTCC CAGACGCAGG	ACGGTAAAGA TGCCATTTCT	CCCCGGGCAG
5251					TCTAGCGCCT AGATCGCGGA
5301					GAGTGGGGGA CTCACCCCT
5,351	CCCCATGGCA GGGGTACCGT	TGGGGTGGGT	CTCGCGCCTC	GCGTACATGC CGCATGTACG	CGCAAATGTC GCGTTTACAG
5401	GTAAACGTAG CATTTGCATC	AGGGGCTCTC TCCCCGAGAG	TGAGTATTCC ACTCATAAGG	AAGATATGTA TTCTATACAT	GGGTAGCATC CCCATCGTAG
5451	TTCCACCGCG AAGGTGGCGC	GATGCTGGCG CTACGACCGC	GCGTGCATTA	CGTATAGTTO	GTGCGAGGGA GCACGCTCCCT
5501	GCGAGGAGGT CGCTCCTCCA	CGGGACCGAG GCCCTGGCTC	GTTGCTACGC CAACGATGCC	GCCCGACGACGACGACGACGACGACGACGACGACGACGAC	CTGCTCGGAA A GACGAGCCTT
5551	GACTATCTGC CTGATAGACC	CTGAAGATGO GACTTCTACO	CATGTGAGTT GTACACTCA	GGATGATATO A CCTACTATAC	G GTTGGACGCT C CAACCTGCGA

Figure 27F

5651				AGCTCGGCGG TCGAGCCGCC	
5701				GATGATGTCA CTACTACAGT	
5751				GGACAAACTC CCTGTTTGAG	
5801				GCCTCCGAAC CGGAGGCTTG	
5851				GGCGCAGCAT CCGCGTCGTA	
5901				GGAGCGAGGT CCTCGCTCCA	
5951				TACTGGTATT ATGACCATAA	
6001				AAAGTCCGTG TTTCAGGCAC	
6051				CGTTGAAGAG GCAACTTCTC	
6101				AAGGGTCCCG TTCCCAGGGC	
6151				GATCTCGTCA CTAGAGCAGT	
6201				AGCGCGGGAT TCGCGCCCTA	
6251	-			AGCTCTTCAG TCGAGAAGTC	
6301					GAAGCGACGA CTTCGCTGCT
6351	ATGAGCTCCA TACTCGAGGT	CAGGTCACGG GTCCAGTGCC	GCCATTAGCA CGGTAATCGT	TTTGCAGGTG AAACGTCCAC	GTCGCGAAAG CAGCGCTTTC
6401					TGCAGTAGAA ACGTCATCTT
6451					GCGGCTAGGT CGCCGATCCA
6501					CATGACCAGC GTACTGGTCG

Figure 27G

6601	TACATCGTAG ATGTAGCATC				
6651	GGAAGAACTG CCTTCTTGAC		CACCAATTGG GTGGTTAACC		
6701	TGAAAGTAGA ACTTTCATCT		ACGGGCCGAA TGCCCGGCTT		
6751			AGCGGTGCAC TCGCCACGTG		
6801 .	GGTTGACCTG CCAACTGGAC		ACAAGGAAGC TGTTCCTTCG		
6851			GTGGTCTTCT CACCAGAAGA		
6901	ACCGTCTGGC TGGCAGACCG		GAGTTACGGT CTCAATGCCA		
6951			TCCGCGCGCG AGGCGCGCGC		
7001			GTCCATGGTC CAGGTACCAG		
7051			GGTTTACCTC CCAAATGGAG		
7101			CTAATTTCCA GATTAAAGGT		
7151			GCATCCCCGC CGTAGGGGCG		
7201			GGGTGTCCTT CCCACAGGAA		
7251	GTGACGCGGG CACTGCGCCC	CGAGCCCCCG	GAGGTAGGGG CTCCATCCCC	GGGCTCCGGA CCCGAGGCCT	CCCGCCGGGA GGGCGGCCCT
7301	GAGGGGGCAG CTCCCCCGTC	GGGCACGTCG CCCGTGCAGC	CCCCCCCCC	CGGGCAGGAG	CTGGTGCTGC GACCACGACG
7351	GCGCGTAGGT CGCGCATCCA	TGCTGGCGAA ACGACCGCTT	CGCGACGACG CGCCTGCTGC	CGGCGGTTGA GCCGCCAACT	TCTCCTGAAT AGAGGACTTA
7401	CTGGCGCCTC GACCGCGGAG	TGCGTGAAGA ACGCACTTCT	CGACGGGCCC	GGTGAGCTTG CCACTCGAAC	AACCTGAAAG TTGGACTTTC
7451	AGAGTTCGAC TCTCAAGCTG	AGAATCAATT	TCGGTGTCGT AGCCACAGCA	TGACGGCGGC	CTGGCGCAAA GACCGCGTTT

Figure 27H

7551		TCTTCCTCCT AGAAGGAGGA			
7601		GTCGTTGGAA CAGCAACCTT			
7651		CGTTCCAGAC GCAAGGTCTG			
7701		ATGACCACCT TACTGGTGGA			
7751		GTTTCGCAGG CAAAGCGTCC			
7801		CCACGAAGAA GGTGCTTCTT			
7851		CCCAAGGCCT GGGTTCCGGA			
7901	CGGCGAAGTT GCCGCTTCAA	GAAAAACTGG CTTTTTGACC	GAGTTGCGCG CTCAACGCGC	CCGACACGGT GGCTGTGCCA	TAACTCCTCC ATTGAGGAGG
7951		GGATGAGCTC CCTACTCGAG			CGCGCTCAAA GCGCGAGTTT
8001					ATAAGGGCCT TATTCCCGGA
8051					ACGGCGCGA .TGCCGCCGCT
8101					TCTCCCGCG AGAGGGGCGC
8151					CGGGGGCGCA
8201	GTTGGAAGAC CAACCTTCTG	GCCGCCCGTC	ATGTCCCGGT TACAGGGCCA	TATGGGTTGG ATACCCAACC	CGGGGGGCTG CGCCCCCGAC
8251	CCATGCGGCA GGTACGCCGT	GGGATACGGC	GCTAACGATG CGATTGCTAC	CATCTCAACA GTAGAGTTGT	ATTGTTGTGT TAACAACACA
8301	AGGTACTCCG TCCATGAGGC	CCGCCGAGGG	ACCTGAGCGA TGGACTCGCT	GTCCGCATCG CAGGCGTAGC	ACCGGATCGG TGGCCTAGCC
8351	AAAACCTCTC TTTTGGAGAG	GAGAAAGGCG CTCTTTCCGC	TCTAACCAGT AGATTGGTCA	CACAGTCGCA GTGTCAGCGT	A AGGTAGGCTG TCCATCCGAC
8401	AGCACCGTGG TCGTGGCACC	GCCCGCCGTC	: GCCCGCCGCG	TCGGGGTTGT	TTCTGGCGGA A AAGACCGCCT

Figure 27I

•					
8501		CACCATGTCC GTGGTACAGG			
8551	TCGGCCATGC AGCCGGTACG	CCCAGGCTTC GGGTCCGAAG	GTTTTGACAT CAAAACTGTA	CGGCGCAGGT GCCGCGTCCA	CTTTGTAGTA GAAACATCAT
8601	GTCTTGCATG CAGAACGTAC	AGCCTTTCTA TCGGAAAGAT	CCGGCACTTC GGCCGTGAAG	TTCTTCTCCT AAGAAGAGGA	TCCTCTTGTC AGGAGAACAG
8651	CTGCATCTCT GACGTAGAGA	TGCATCTATC ACGTAGATAG	GCTGCGGCGG CGACGCCGCC	CGGCGGAGTT GCCGCCTCAA	TGGCCGTAGG ACCGGCATCC
8701	TGGCGCCCTC ACCGCGGGAG	TTCCTCCCAT AAGGAGGGTA	GCGTGTGACC CGCACACTGG	CCGAAGCCCC GGCTTCGGGG	TCATCGGCTG AGTAGCCGAC
8751	AAGCAGGGCT TTCGTCCCGA	AGGTCGGCGA TCCAGCCGCT	CAACGCGCTC GTTGCGCGAG	GGCTAATATG CCGATTATAC	GCCTGCTGCA CGGACGACGT
8801	CCTGCGTGAG GGACGCACTC	GGTAGACTGG CCATCTGACC	AAGTCATCCA TTCAGTAGGT	TGTCCACAAA ACAGGTGTTT	GCGGTGGTAT CGCCACCATA
8851	GCGCCCGTGT CGCGGGCACA	TGATGGTGTA ACTACCACAT	AGTGCAGTTG TCACGTCAAC	GCCATAACGG CGGTATTGCC	ACCAGTTAAC TGGTCAATTG
8901	GGTCTGGTGA CCAGACCACT	CCCGGCTGCG GGGCCGACGC	AGAGCTCGGT TCTCGAGCCA	GTACCTGAGA CATGGACTCT	CGCGAGTAAG GCGCTCATTC
8951	CCCTCGAGTC GGGAGCTCAG	AAATACGTAG TTTATGCATC	TCGTTGCAAG AGCAACGTTC	TCCGCACCAG AGGCGTGGTC	GTACTGGTAT CATGACCATA
9001	CCCACCAAAA	AGTGCGGCGG	CGGCTGGCGG	TAGAGGGGCC	AGCGTAGGGT
9051	GGCCGGGCT	CCGGGGGCGA	GATCTTCCAA CTAGAAGGTT	CATAAGGCGA	TGATATCCGT ACTATAGGCA
9101	AGATGTACCI	GGACATCCAG	GTGATGCCGG	CGGCGGTGGT	GGAGGCGCGC
9151	GGAAAGTCGC	GGACGCGGTT CCTGCGCCAA	CCAGATGTTG	CGCAGCGGCA	AAAAGTGCTC
9201	CATGGTCGG	ACGCTCTGGC	CGGTCAGGCG	CGCGCAATCG	TTGACGCTCT AACTGCGAGA
9251	AGACCGTGC	AAAGGAGAGC	CTGTAAGCG	GCACTCTTCC	GTGGTCTGGT
9301	GGATAAATTO	GCAAGGGTAT	CATGGCGGA	GACCGGGGTT	CACCAGACCA
9351					GCTCGGGGCA GTGTCGAACCC
,,,,	TAGGCCGGC	A GGCGGCACTA	GGTACGCCA	A TGGCGGGCG	ACAGCTTGGG

Figure 27J

9451		CTGCTGCGCT GACGACGCGA		
9501	TAAGCGGTTA ATTCGCCAAT	GGCTGGAAAG CCGACCTTTC		
9551	CCGGAGGGTT GGCCTCCCAA	ATTTTCCAAG TAAAAGGTTC		
9601		CGGACTGCGG GCCTGACGCC		
9651	GACCCCGCTT CTGGGGCGAA	GCAAATTCCT CGTTTAAGGA		
9701		GCATCCGGTG CGTAGGCCAC	•	
9751	••••	AAGAGCAGCG TTCTCGTCGC		
9801		GGAGGGGCGA CCTCCCGCT		
9851		CCCGCGGCGC		
9901		TGGCGCGGCT ACCGCGCCGA		
9951		AAGCGTGATA TTCGCACTAT		
10001		CCGCGAGGGA GGCGCTCCCT		
10051				AGCGGTTGCT TCGCCAACGA
10101	GCGCGAGGAG CGCGCTCCTC			AGTCCCGCGC TCAGGGCGCG
10151	GCGCACACGT CGCGTGTGCA			GCAGACGGTG CGTCTGCCAC
10201	AACCAGGAGA TTGGTCCTCT			TGCGTACGCT ACGCATGCGA
10251	TGTGGCGCGC ACACCGCGCG			TGGGACTTTG ACCCTGAAAC
10301	TAAGCGCGCT ATTCGCGCGA			GGCGCAGCTG CCGCGTCGAC

Figure 27K

10401	GCTAAACATA	GTAGAGCCCG	AGGGCCGCTG	GCTGCTCGAT	TTGATAAACA
	CGATTTGTAT	CATCTCGGGC	TCCCGGCGAC	CGACGAGCTA	AACTATTTGT
10451				GCTTGAGCCT CGAAC,TCGGA	
10501				CTGGGCAAGT GACCCGTTCA	
10551	CAAGATATAC	CATACCCCTT	ACGTTCCCAT	AGACAAGGAG	GTAAAGATCG
	GTTCTATATG	GTATGGGGAA	TGCAAGGGTA	TCTGTTCCTC	CATTTCTAGC
10601				TGCTTACCTT ACGAATGGAA	
10651	CTGGGCGTTT	ATCGCAACGA	GCGCATCCAC	AAGGCCGTGA	GCGTGAGCCG
	GACCCGCAAA	TAGCGTTGCT	CGCGTAGGTG	TTCCGGCACT	CGCACTCGGC
10701	GCGGCGCGAG	CTCAGCGACC	GCGAGCTGAT	GCACAGCCTG	CAAAGGGCCC
	CGCCGCGCTC	GAGTCGCTGG	CGCTCGACTA	CGTGTCGGAC	GTTTCCCGGG
10751	TGGCTGGCAC	GGGCAGCGGC	GATAGAGAGG	CCGAGTCCTA	CTTTGACGCG
	ACCGACCGTG	CCCGTCGCCG	CTATCTCTCC	GGCTCAGGAT	GAAACTGCGC
10801	GGCGCTGACC	TGCGCTGGGC	CCCAAGCCGA	CGCGCCCTGG	AGGCAGCTGG
	CCGCGACTGG	ACGCGACCCG	GGGTTCGGCT	GCGCGGGACC	TCCGTCGACC
10851	GGCCGGACCT	GGGCTGGCGG	TGGCACCCGC	GCGCGCTGGC	AACGTCGGCG
	CCGGCCTGGA	CCCGACCGCC	ACCGTGGGCG	CGCGCGACCG	TTGCAGCCGC
10901	GCGTGGAGGA	ATATGACGAG	GACGATGAGT	ACGAGCCAGA	GGACGGCGAG
	CGCACCTCCT	TATACTGCTC	CTGCTACTCA	TGCTCGGTCT	CCTGCCGCTC
10951	TACTAAGCGG ATGATTCGCC	TGATGTTTCT	GATCAGATGA CTAGTCTACT	TGCAAGACGC ACGTTCTGCG	AACGGACCCG TTGCCTGGGC
11001	GCGGTGCGGG	CGGCGCTGCA	GAGCCAGCCG	TCCGGCCTTA	ACTCCACGGA
	CGCCACGCCC	CGCGCGACGT	CTCGGTCGGC	AGGCCGGAAT	TGAGGTGCCT
11051	CGACTGGCGC	CAGGTCATGG	ACCGCATCAT	GTCGCTGACT	GCGCGCAATC
	GCTGACCGCG	GTCCAGTACG	TGGCGTAGTA	CAGCGACTGA	CGCGCGTTAG
11101	CTGACGCGTT GACTGCGCAA	CCGGCAGCAC GGCCGTCGTC	CCGCAGGCCA CGCGTCCGGT	A ACCGGCTCTC TGGCCGAGAG	CGCAATTCTG GCGTTAAGAC
11151	GAAGCGGTGG CTTCGCCACG	TCCCGGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCGC	GCGTTTGGGC	ACGCACGAGA TGCGTGCTC1	A AGGTGCTGGC TCCACGACCG
11201	GATCGTAAAC CTAGCATTTC	GCGCTGGCCGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	AAAACAGGGGCCCCCCCCCCCCCCCCCCCCCCCCCCCC	CATCCGGCCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	GACGAGGCCG CTGCTCCGGC
11251	GCCTGGTCTA CGGACCAGA	A CGACGCGCTO CGCGCGCGA	G CTTCAGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCGCG	TGGCTCGTTA	A CAACAGCGGC T GTTGTCGCCG

Figure 27L

11351			CAACCTGGGC GTTGGACCCG	
11401			CCAACGTGCC GGTTGCACGG	
11451			CGGCTAATGG GCCGATTACC	
11501			AGACTATTTT TCTGATAAAA	
11551			GCCAGGCTTT CGGTCCGAAA	
11601			GGCGACCGCG CCGCTGGCGC	
13.651			GCTGCTGCTA CGACGACGAT	
11701			CATACCTAGG GTATGGATCC	
11751			CATGTGGACG GTACACCTGC	
11801			GGGGCAGGAG CCCCGTCCTC	
11851			CCAACCGGCG GGTTGGCCGC	
11901			GAGCGCATTT CTCGCGTAAA	
11951			CGACGGGGTA GCTGCCCCAT	
12001	TGGCGCTGGA ACCGCGACCT		AACCGGGCAT TTGGCCCGTA	
12051	AACCGGCCGT TTGGCCGGCA		TACTTGCATC ATGAACGTAG	
12101	CGTGAACCCC GCACTTGGGG			CACTGGCTAC GTGACCGATG
12151	CGCCCCCTGG GCGGGGGACC			GGGTAACGAT CCCATTGCTA
12201	GGATTCCTCT CCTAAGGAGA			CGCAACCGCA GCGTTGGCGT

Figure 27 M

12301			AGCAGCTTGT TCGTCGAACA		
12351			CCCATTTCCA GGGTAAAGGT		
12401			CGCGCCTGCT GCGCGGACGA		
12451			CAGCGCGAAA GTCGCGCTTT		
12501			CCTAGTGGAC GGATCACCTG		
12551	CATGCGCGTC	CTCGTGTCCC	ACGTGCCAGG TGCACGGTCC	GGCCCGGGC	GGGTGGGCAG
12601			CGGGGTCTGG GCCCCAGACC		
12651			GGATTTGGGA CCTAAACCCT		
12701			GGAGAATGTT CCTCTTACAA		
12751	TACGTTTTAT	TTTTTGAGTG	GTTCCGGTAC	CGTGGCTCGC	
12801	ACATAAGGGG	AATCATACGC	CGCGCGCCGC	TACATACTCC	AAGGTCCTCC TTCCAGGAGG
12851	AGGGAGGATG	CTCTCACACC	ACTCGCGCCG	CGGTCACCGC	GCGGCGCTGG
12901					TCCGCGGTAC AGGCGCCATG
12951	CTGCGGCCTA GACGCCGGAT	CCGGGGGGAG	AAACAGCATC TTTGTCGTAG	CGTTACTCTG GCAATGAGAC	AGTTGGCACC TCAACCGTGG
13001					TCAACGGATG AGTTGCCTAC
13051					GACCACGGTC CTGGTGCCAG
13101					AGACCATCAA TCTGGTAGTT
13151					ATCCTGCATA TAGGACGTAT

Figure 27N

13251		TGTCGCGCTT ACAGCGCGAA			
13301	. =	GTGGAGTTCA CACCTCAAGT		-	
13351		CCTTATGAAC GGAATACTTG			
13401		ACGGGGTTCT TGCCCCAAGA			
13451		AGACTGGGGT TCTGACCCCA			
13501		AAACGAAGCC TTTGCTTCGG			
13551		ACTTCACCCA TGAAGTGGGT			
13601		CCCTTCCAGG GGGAAGGTCC			
13651		CATTCCCGCA GTAAGGGCGT			
13701	• • • • • • • •	ACACCGAACA TGTGGCTTGT			
13751	•	GGCGCGGAAG CCGCGCCTTC			
13801		GGACATGAAC CCTGTACTTG			
13851		AGGAGAAGCG TCCTCTTCGC	•		
13901	CGCCCCCGCT GCGGGGGCGA	GCGCAACCCG CGCGTTGGGC			
13951		GACAGAGGAC CTGTCTCCTG			CCTAATAAGC GGATTATTCG
14001	AATGACAGCA TTACTGTCGT	CCTTCACCCA GGAAGTGGGT			
14051	CGGCGACCCT GCCGCTGGGA	CAGACCGGAA GTCTGGCCTT			
14101	ACGTAACCTG TGCATTGGAC	CGGCTCGGAG GCCGAGCCTC			

Tigure 270

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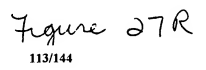
14201		GAGCTGTTGC CTCGACAACG		_	
14251		CTCCCAACTC GAGGGTTGAG			
14301		TTCCCGAGAA AAGGGCTCTT	· · - · · - · · · ·		
14351		GTCAGTGAAA CAGTCACTTT			
14401		CAACAGCATC GTTGTCGTAG			
14451		GCACCTGCCC CGTGGACGGG			
14501	- -	CTATCGAGCC GATAGCTCGG		· - - ·	
14551		CAATAACACA GTTATTGTGT			
14601		CCAAGAAGCG GGTTCTTCGC			
14651		GCGCCCTGGG CGCGGGACCC			
14701		TGACGCCATC ACTGCGGTAG			
14751		CGCCACCAGT GCGGTGGTCA			
14801	• •	GCCCGGCGCT CGGGCCGCGA			
14851	TAGCACGTCG ATCGTGCAGC	CCACCGCCGC GGTGGCGGCG			
14901	GCGGCCCTGC CGCCGGGACG	TTAACCGCGC AATTGGCGCG			
14951	GGCCGCTCGA CCGGCGAGCT	AGGCTGGCCG TCCGACCGGC			
15001	GGCGACGAGC CCGCTGCTCG	GGCCGCCGCA CCGGCGGCGT		=	
15051	GGTCGCAGGG CCAGCGTCCC	GCAACGTGTA CGTTGCACAT			

FIGURE 27P

15151				GCCGCCGCCG CGGCGGCGGC	
15201				ATGCTCCAGG TACGAGGTCC	
15251	GGAGATCTAT CCTCTAGATA			GCAGGATTAC CGTCCTAATG	
15301				ATGATGATGA TACTACTACT	
15351				CCCAGGCGAC GGGTCCGCTG	
15401				ACCCGGCACC TGGGCCGTGG	
15451	TTACGCCCGG AATGCGGGCC	TGAGCGCTCC ACTCGCGAGG	ACCCGCACCT TGGGCGTGGA	ACAAGCGCGT TGTTCGCGCA	GTATGATGAG CATACTACTC
15501	GTGTACGGCG CACATGCCGC	ACGAGGACCT TGCTCCTGGA	GCTTGAGCAG CGAACTCGTC	GCCAACGAGC CGGTTGCTCG	GCCTCGGGGA CGGAGCCCCT
15551				GCTGGCGTTG CGACCGCAAC	
15601				TAACACTGCA ATTGTGACGT	
15651				GGCCTAAAGC CCGGATTTCG	GCGAGTCTGG CGCTCAGACC
15701					CAGCGACTGG GTCGCTGACC
15751	AAGATGTCTT TTCTACAGAA	GGAAAAAATG	ACCGTGGAAC TGGCACCTTG	CTGGGCTGGA GACCCGACCT	GCCCGAGGTC CGGGCTCCAG
15801	CGCGTGCGGC	CAATCAAGCA GTTAGTTCGT	GGTGGCGCCGC	GGACTGGGCG CCTGACCCGC	TGCAGACCGT ACGTCTGGCA
15851	GGACGTTCAG CCTGCAAGTC	ATACCCACTA TATGGGTGAT	CCAGTAGCAC GGTCATCGTG	CAGTATTGCC GTCATAACGC	ACCGCCACAG TGGCGGTGTC
15901	AGGGCATGGA TCCCGTACCT	GACACAAACO CTGTGTTTGC	TCCCCGGTTG AGGGGCCAAC	CCTCAGCGGT GGAGTCGCCA	GGCGGATGCC CCGCCTACGG
15951	GCGGTGCAGG CGCCACGTCC	CGGTCGCTGC CCAGCGACC	GGCCGCGTCC GCCGCGCAGC	AAGACCTCTA TTCTGGAGA1	CGGAGGTGCA CGCCTCCACGT
16001	AACGGACCCG TTGCCTGGGC	TGGATGTTTC ACCTACAAA	GCGTTTCAGC GCGCAAAGTCC	Geeeeccec	CCGCGCCGTT GGCGCGGCAA

Figure 270

16051				TGCCCGAATA ACGGGCTTAT	
16101				GGCTACACCT CCGATGTGGA	
16151				CACTGGAACC GTGACCTTGG	
16201				TTTCCGTGCG AAAGGCACGC	
16251				ACAGCGCGCT TGTCGCGCGA	
16301				TGCAGATATG ACGTCTATAC	
16351				GAGGAAGAAT CTCCTTCTTA	
16401	AGGGGCATGG TCCCCGTACC			GGCATGCGTC CCGTACGCAG	
16451				GCGCGGCGGT CGCGCCGCCA	
16501				GCGCCGTGCC CGCGCACGG	
16551				TTAAAAACAA AATTTTTGTT	GTTGCATGTG CAACGTACAC
16601					GGTCCTGTAA CCAGGACATT
16651	GATAAAACAT	CTTACCTTCT	GTAGTTGAAA	CGCAGAGACC	CCCCGCGACA
	GCCGAGCGCG	GGCAAGTACC	CTTTGACCGI	TCTATAGCCG	ACCAGCAATA TGGTCGTTAT
16751	ACTCGCCACC	GCGGAAGTCG	ACCCCGAGCG	ACACCTCGCC	CATTAAAAAT GTAATTTTTA
16801	AAGCCAAGGT	GGCAATTCTT	GATACCGTCC	TTCCGGACCT	ACAGCAGCAC TGTCGTCGTG
16851	AGGCCAGAT(TCCGGTCTA(GACTCCCTAT	AGTTGAAAGA TCAACTTTC	A GCAAAATTTO CGTTTTAAAG	CAACAAAAGG GTTGTTTTCC
16901	ACCATCTACO	GGACCGGAGA	CCGTAATCG	CCCACCACCI	A CCTGGCCAAC C GGACCGGTTG
16951					GCCCTCCCGT GCGGAGGGCA



17051		GCGCCCCGAC CGCGGGGCTG			
17161		CGTACGAGGA GCATGCTCCT			
17151		CCCATGGCTA GGGTACCGAT			
17201		GCCTCCCCC CGGAGGGGGG			
17251		CCGTTGTTGT GGCAACAACA			
17301		GGTCCGCGAT CCAGGCGCTA			
17351		GAACAGCATC CTTGTCGTAG			
17401		TCTGATAGCT AGACTATCGA			
17451		CCAGAGGAGC GGTCTCCTCG			
17501		CTTCGATGAT GAAGCTACTA			
17551		TCGGAGTACC AGCCTCATGG			
17601		GTACTTCAGC CATGAAGTCG			
17651	CGCGGATGCG	ACGACGTGAC TGCTGCACTG	GTGTCTGGCC	AGGGTCGCAA	ACTGCGACGC
		CACCTGGCAC	TCCTATGACG	CATGAGCATG	TTCCGCGCCA
		ACACCCACTA	TTGGCACACG	ACCTGTACCG	AAGGTGCATG
		CGCCGCACGA	CCTGTCCCCG	GGATGAAAAT	TCGGGATGAG
		ATGTTGCGGG	ACCGAGGGTT	CCCACGGGGT	TTAGGAACGC
17901	AATGGGATGA TTACCCTACT	AGCTGCTACT TCGACGATGA			

Figure 275

17951	GATGACAACG CTACTGTTGC				
18001	CGTATTTGGG GCATAAACCC				
18051	TTCAAATAGG AAGTTTATCC	TGTCGAAGGT ACAGCTTCCA			
18101		CTCAAATAGG GAGTTTATCC			
18151		GGGAGAGTCC CCCTCTCAGG			
18201	TGCCAAGTAT	TGCAAAACCC ACGTTTTGGG	TGTTTACTTT	TACCTCCCGT	TCCGTAAGAA
18251	GTAAAGCAAC CATTTCGTTG	AAAATGGAAA TTTTACCTTT	GCTAGAAAGT CGATCTTTCA	CAAGTGGAAA GTTCACCTTT	TGCAATTTTT ACGTTAAAAA
18301	CTCAACTACT GAGTTGATGA	GAGGCAGCCG CTCCGTCGGC	CAGGCAATGG GTCCGTTACC	TGATAACTTG ACTATTGAAC	ACTCCTAAAG TGAGGATTTC
18351		CAGTGAAGAT GTCACTTCTA			CACTCATATT GTGAGTATAA
18401	TCTTACATGC AGAATGTACG	CCACTATTAA GGTGATAATT	GGAAGGTAAC CCTTCCATTG	TCACGAGAAC AGTGCTCTTG	TAATGGGCCA ATTACCCGGT
18451	ACAATCTATG TGTTAGATAC	CCCAACAGGC GGGTTGTCCG	CTAATTACAT GATTAATGTA	TGCTTTTAGG ACGAAAATCC	GACAATTTTA CTGTTAAAAT
18501	TTGGTCTAAT AACCAGATTA	GTATTACAAC CATAATGTTG	AGCACGGGTA TCGTGCCCAT	ATATGGGTGT TATACCCACA	TCTGGCGGGC AGACCGCCCG
18551	CAAGCATCGC GTTCGTAGCG	AGTTGAATGC TCAACTTACG	TGTTGTAGAT ACAACATCTA	TTGCAAGACA AACGTTCTGT	GAAACACAGA CTTTGTGTCT
18601	GCTTTCATAC CGAAAGTATG	CAGCTTTTGC GTCGAAAACG	TTGATTCCAT AACTAAGGTA	TGGTGATAGA ACCACTATCT	ACCÁGGTACT TGGTCCATGA
	AAAGATACAC	CTTAGTCCGA	CAACTGTCGA	TACTAGGTCT	TGTTAGAATT ACAATCTTAA
	TAACTTTTAG	TACCTTGACT	TCTACTTGAA	GGTTTAATGA	GCTTTCCACT CGAAAGGTGA
18751	GGGAGGTGT(CCCTCCACA(ATTAATACAC TAATTATGTC	AGACTCTTAC TCTGAGAATC	CAAGGTAAAA GTTCCATTT	CCTAAAACAG GGATTTTGTC
18801	GTCAGGAAA/ CAGTCCTTT	A TGGATGGGAL T ACCTACCCT	A AAAGATGCTA TTTCTACGAT	CAGAATTTT(GTCTTAAAA(AGATAAAAAT TCTATTTTA
18851	GAAATAAGA(CTTTATTCT(TTGGAAATA	A TTTTGCCATO	GAAATCAATO CTTTAGTTA	TAAATGCCAA ATTTACGGTT

Figure 27 T

18951		CAGTCCTTCC			
	TCGATTTCAT	GTCAGGAAGG	TIGCATTTT	AAAGACTATT	GGGTTTGTGG
19001	TACGACTACA	TGAACAAGCG	AGTGGTGGCT	CCCGGGCTAG	TGGACTGCTA
	ATGCTGATGT	ACTTGTTCGC	TCACCACCGA	GGGCCCGATC	ACCTGACGAT
19051	••••	GGAGCACGCT			
	GTAATTGGAA	CCTCGTGCGA	CCAGGGAACT	GATATACCTG	TTGCAGTTGG
19101	CATTTAACCA	CCACCGCAAT	GCTGGCCTGC	GCTACCGCTC	AATGTTGCTG
	GTAAATTGGT	GGTGGCGTTA	CGACCGGACG	CGATGGCGAG	TTACAACGAC
19151	CCCLATGGTC	GCTATGTGCC	CTTCCACATC	CAGGTGCCTC	AGAAGTTCTT
17171		CGATACACGG			
19201		AACCTCCTTC			
	ACGGTAATTT	TTGGAGGAAG	AGGACGGCCC	GAGTATGTGG	ATGCTCACCT
19251	ACTTCAGGAA	GGATGTTAAC	ATGGTTCTGC	AGAGCTCCCT	AGGAAATGAC
	TGAAGTCCTT	CCTACAATTG	TACCAAGACG	TCTCGAGGGA	TCCTTTACTG
19301	C#3 3 CCC##C	ACGGAGCCAG	ርልጥጥል አርጥጥጥ	CATACCATTT	CCCTTTACCC
19301		TGCCTCGGTC			
	GAT TUUCAAU	TGCCTCGGTC	GIAATICAAA		CGGAAA1GCG
19351	CACCTTCTTC	CCCATGGCCC	ACAACACCGC	CTCCACGCTT	GAGGCCATGC
	GTGGAAGAAG	GGGTACCGGG	TGTTGTGGCG	GAGGTGCGAA	CTCCGGTACG
19401		CACCAAÇGAC			
	AATCTTTGCT	GTGGTTGCTG	GTCAGGAAAT	TGCTGÄTAGA	GAGGCGGCGG
19451		ACCCTATACC			
	TTGTACGAGA	TGGGATATGG	GCGGTTGCGA	TGGTTGCACG	GGTATAGGTA
				amaaaaaamma	3.000000mm3
19501		AACTGGGCGG			
	GGGGAGGGCG	TTGACCCGCC	GAAAGGCGCC	GACCCGGAAG	TGCGCGGAAT
19551		AACCCCATCA			
	TCTGATTCCT	TTGGGGTAGT	GACCCGAGCC	CGATGCTGGG	AATAATGTGG
19601	TACTCTGGCT	СТАТАСССТА	CCTAGATGGA	ACCTTTTACC	TCAACCACAC
19001					AGTTGGTGTG
	AIGAGACCGA	GAIAIGGGAI	GGATCTACCT	1003391100	
19651	CTTTAAGAAG	GTGGCCATTA	CCTTTGACTC	TTCTGTCAGC	TGGCCTGGCA
	GAAATTCTTC	CACCGGTAAT	GGAAACTGAG	AAGACAGTCG	ACCGGACCGT
19701	ATGACCGCCT	GCTTACCCCC	AACGAGTTTG	AAATTAAGCG	CTCAGTTGAC
17,01					GAGTCAACTG
19751					ACTGGTTCCT
	CCCCTCCCAA	TGTTGCAACG	GGTCACATTG	TACTGGTTTC	TGACCAAGGA
				CD1001000	
19801					TTCTATATCC
	CCATGTTTAC	GATCGATTGA	TATTGTAACC	GATGGTCCCG	AAGATATAGG

Figure 274

19851			ATGTACTCCT TACATGAGGA		
19901			TGATACTAAA ACTATGATTT		
19951			ACAACTCTGG TGTTGAGACC		
20001			GCCTACCCTG CGGATGGGAC		
20051			CAGCATTACC GTCGTAATGG		
20101			CATTCTCCAG GTAAGAGGTC		
20151			AACCTTCTCT TTGGAAGAGA		
20201			GGATCCCATG CCTAGGGTAC		
20251			ACGTGGTCCG TGCACCAGGC		GGCGTGGCGC CCGCACCGCG
20301			CTGCGCACGC GACGCGTGCG		
20351					CATGGGCTCC GTACCCGAGG
20401					GTGGGCCATA CACCCGGTAT
20451					TCTCCACACA AGAGGTGTGT
20501	AGCTCGCCTG TCGAGCGGAC	CGCCATAGTC GCGGTATCAG	AATACGGCCG TTATGCCGGC	GTCGCGAGAC CAGCGCTCTG	TGGGGGCGTA
20551					GCTACCTCTT CGATGGAGAA
20601					TACCAGTTTG ATGGTCAAAC
20651	AGTACGAGTC TCATGCTCAG	ACTCCTGCGC TGAGGACGCG	CGTAGCGCCA GCATCGCGGT	TTGCTTCTTC	CCCCGACCGC GGGGCTGGCG
20701					CCAACTCGGC GGTTGAGCCG
20751					GCCAACTGGC CGGTTGACCG

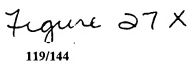
Figure 27 V.

20851	CCCAACTCCA GGGTTGAGGT			CAGCCCACCC GTCGGGTGGG	
20901	GGTCCTTGTC	GAGATGTCGA	AGGACCTCGC	CCACTCGCCC GGTGAGCGGG	ATGAAGGCGT
20951	CGGTGTCACG	CGTCTAATCC	TCGCGGTGAA	CTTTTTGTCA GAAAAACAGT	GAACTTTTTG
21001	TACATTTTTA	TTACATGATC	TCTGTGAAAG	AATAAAGGCA TTATTTCCGT	TTACGAAAAT
21051	AAACATGTGA	GAGCCCACTA	ATAAATGGGG	CACCCTTGCC GTGGGAACGG	CAGACGCGGC
21101	AAATTTTTAG	TTTCCCCAAG	ACGGCGCGTA	CGCTATGCGC GCGATACGCG	GTGACCGTCC
21151	CTGTGCAACG	CTATGACCAC	AAATCACGAG	CACTTAAACT GTGAATTTGA	GTCCGTGTTG
21201	GTAGGCGCCG	TCGAGCCACT	TCAAAAGTGA	CCACAGGCTG GGTGTCCGAC	GCGTGGTAGT
21251	GGTTGCGCAA	ATCGTCCAGC	CCGCGGCTAT	AGAACTTCAG	
21301	GGAGGCGGGA	CGCGCGCGCT	CAACGCTATG	TGTCCCAACG	AGCACTGGAA TCGTGACCTT
21351	GTGATAGTCG	CGGCCCACCA	CGTGCGACCG	GTCGTGCGAG	TTGTCGGAGA AACAGCCTCT
21401	AGTCTAGGCG	CAGGTCCAGG	AGGCGCAACG	AGTCCCGCTT	CGGAGTCAAC
21451	AAACCATCGA	CGGAAGGGTT	TTTCCCGCGC	: ACGGGTCCGA	TTGAGTTGCA AACTCAACGT
	GAGCGTGGCA	TCACCGTAG	TTTCCACTGO	CACGGGCCAG	TGGGCGTTAG ACCCGCAATC
	CTATGTCGCC	GACGTATTT	CGGAACTAGA	A CGAATTTTC	CACCTGAGCC GTGGACTCGG
	AAACGCGGA	GTCTCTTCTT	r GTACGGCGT	r ctgaacggc	AAAACTGATT TTTTGACTAA
	CCGGCCTGTC	CGGCGCAGC	A CGTGCGTCG	r GGAACGCAG	G GTGTTGGAGA C CACAACCTCT
21701	TCTGCACCA(AGACGTGGT(ATTTCGGCCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	CACCGGTTC	TCACGATCT	r GGCCTTGCTA A CCGGAACGAT

7, gure 27 W

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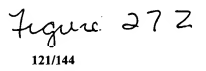
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21851			CGGTGCAGCC GCCACGTCGG	= -	
21901		· · · · · · · · · · ·	CTCTGCAAAC GAGACGTTTG		
21951		· · · · · · · · · · · · · · · · · · ·	CAAAGGTCTT GTTTCCAGAA		
22001			TTCAGCCAGG AAGTCGGTCC		
22051			TAGTTTGAAG ATCAAACTTC		•
22101			CGCGCGCGC	-	
22151			CTCAGCGGGT GAGTCGCCCA		
22201			CTCTTCCTCT GAGAAGGAGA		
22251			GCCGCCGCAC CGGCGGCGTG		
22301			GGGTTGCTGA CCCAACGACT		
22351			GCTGTCCACG CGACAGGTGC		
22401			GGCGCTTCTT CCGCGAAGAA		
22451	CCAAATCCGC GGTTTAGGCG				GCGCGGCACC CGCGCCGTGG
22501	AGCGCGTCTT TCGCGCAGAA				TACGCCGCCT ATGCGGCGGA
22551	CATCCGCTTT GTAGGCGAAA		CCCGGGGAGG GGGCCCCTCC		
22601	ACGACACGTC TGCTGTGCAG				GCGTCCGCGC CGCAGGCGCG
22651	TCGGGGGTGG AGCCCCCACC				TTTCCTTCTC AAAGGAAGAG



22751		TGAGTTCGCC ACTCAAGCGG		
22801		TCCCCGTCGA AGGGGCAGCT		
22851		GACCCAGGTT CTGGGTCCAA		
22901		GGATAAAAAG CCTATTTTTC		
22951		GGCGGGGGA CCGCCCCCT		
23001		CTGTTGAAGC GACAACTTCG		
23051		AGAGCGCAGC TCTCGCGTCG		
23101		AACGCCACCT TTGCGGTGGA		
23151		ACATGCGAGC TGTACGCTCG		
23201		AGAGGTGCTT TCTCCACGAA		
23251		TATCCTGCCG ATAGGACGGC		
23301		CAGGGCGCTG GTCCCGCGAC		
23351		CTTTGAGGGT GAAACTCCCA		
23401				GAGTGTTGGT CTCACAACCA
23451	GGAACTCGAG CCTTGAGCTC	GGTGACAACG CCACTGTTGC		
23501		CTTTGCCTAC GAAACGGATG		CAAGGTCATG GTTCCAGTAC
23551				CCCTGGAGAG GGGACCTCTC
23601				GCAGTTGGCG CGTCAACCGC

Figure 27 Y

23701	GAGCGACGCA CTCGCTGCGT	AACTAATGAT TTGATTACTA	GGCCGCAGTG CCGGCGTCAC	CTCGTTACCG GAGCAATGGC	TGGAGCTTGA ACCTCGAACT
23751	GTGCATGCAG CACGTACGTC	CGGTTCTTTG GCCAAGAAAC	CTGACCCGGA GACTGGGCCT	GATGCAGCGC CTACGTCGCG	AAGCTAGAGG TTCGATCTCC
23801	AAACATTGCA TTTGTAACGT	CTACACCTTT GATGTGGAAA	CGACAGGGCT GCTGTCCCGA	ACGTACGCCA TGCATGCGGT	GGCCTGCAAG CCGGACGTTC
23851	ATCTCCAACG TAGAGGTTGC	TGGAGCTCTG ACCTCGAGAC	CAACCTGGTC GTTGGACCAG	TCCTACCTTG AGGATGGAAC	GAATTTTGCA CTTAAAACGT
23901			ACGTGCTTCA TGCACGAAGT		
23951	CGCGCCGCGA GCGCGGCGCT	CTACGTCCGC GATGCAGGCG	GACTGCGTTT CTGACGCAAA	ACTTATTTCT TGAATAAAGA	ATGCTACACC TACGATGTGG
24001	TGGCAGACGG ACCGTCTGCC	CCATGGGCGT GGTACCCGCA	TTGGCAGCAG AACCGTCGTC	TGCTTGGAGG ACGAACCTCC	AGTGCAACCT TCACGTTGGA
24051	CAAGGAGCTG GTTCCTCGAC	CAGAAACTGC GTCTTTGACG	TAAAGCAAAA ATTTCGTTTT	CTTGAAGGAC GAACTTCCTG	CTATGGACGG GATACCTGCC
24101	GGAAGTTGCT	CGCGAGGCAC	GCCGCGCACC CGGCGCGTGG	ACCGCCTGTA	GTAAAAGGGG
24151	CTTGCGGACG	AATTTTGGGA	CGTTGTCCCA	GACGGTCTGA	TCACCAGTCA AGTGGTCAGT
24201	TTCGTACAAC	GTCTTGAAAT	CCTTGAAATA	GGATCTCGCG	TCAGGAATCT AGTCCTTAGA
24251	ACGGGCGGTG	GACGACACGT	GAAGGATCGC	TGAAACACGG	CATTAAGTAC GTAATTCATG
24301	GCGCTTACGG	GAGGCGGCGA	AACCCCGGTG	ACGATGGAAG	TGCAGCTAGC ACGTCGATCG
	GTTGATGGAA	CGGATGGTG	GACTGTATTA	A CCTTCTGCAC	AGCGGTGACG TCGCCACTGC
	CAGATGACCT	CACAGTGACA	A GCGACGTTGC	ATACGTGGGG	GCACCGCTCC GCGTGGCGAGG
	GACCAAACGT	TAAGCGTCG	A CGAATTGCT	TCAGTTTAAT	A TCGGTACCTT T AGCCATGGAA
	ACTCGACGTC	CCAGGGAGC	G GACTGCTTT	r CAGGCGCCGI	CCGGGGTTGA GGCCCCAACT
24551	AACTCACTC(TTGAGTGAG(GGGGCTGTGG CCCCGACAC	ACGTCGGCT	r accttcgcal a tggaagcgt	A ATTTGTACCT T TAAACATGGA



24601		ACGCCCACGA TGCGGGTGCT		
24651		GAGCTTACCG CTCGAATGGC	 	-
24701		AGCCATCAAC TCGGTAGTTG		
24751		TTTACTTGGA AAATGAACCT		
24801		CCGCAGCCCT GGCGTCGGGA	 	
24851	AGGATGGCAC TCCTACCGTG	CCAAAAAGAA GGTTTTTCTT		
24901		TGGGACAGTC ACCCTGTCAG		
24951		GGAAGACTGG CCTTCTGACC		
25001		CAGACGAAAC GTCTGCTTTG		
25051		AAATCGGCAA TTTAGCCGTT		
25101		GCCGGCACTG CGGCCGTGAC		
25151		CCAGGGCCGG GGTCCCGGCC		
25201		CAGCGCCAAG GTCGCGGTTC		
25251				CTTCGCCCGC GAAGCGGGCG
25301		TCTACCATCA AGATGGTAGT		ACATCCTGCA TGTAGGACGT
25351		CATCTCTACA GTAGAGATGT		AGCGGCAGCA TCGCCGTCGT
25401				AGACTCTGAC TCTGAGACTG
25451	AAAGCCCAAG TTTCGGGTTC			GGAGCGCTGC CCTCGCGACG
25501	GTCTGGCGCC CAGACCGCGG			AAACAGGATT TTTGTCCTAA

Figure 27 AA

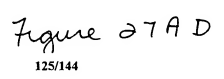
25551	TTTCCCAC	TGTATGCTAT ACATACGATA	ATTTCAACAG TAAAGTTGTC	AGCAGGGG. TCGTCCCCGG	AAGAACAAGA TTCTTGTTCT
25601				CCTCACCCGC GGAGTGGGCG	
25651	ATCACAAAAG TAGTGTTTTC	CGAAGATCAG GCTTCTAGTC	CTTCGGCGCA GAAGCCGCGT	CGCTGGAAGA GCGACCTTCT	CGCGGAGGCT GCGCCTCCGA
25701				AAGGACTAGT TTCCTGATCA	
25751				TCTCCAGCGG AGAGGTCGCC	
25801	CGCCAGCACC GCGGTCGTGG	TGTTGTCAGC ACAACAGTCG	GCCATTATGA CGGTAATACT	GCAAGGAAAT CGTTCCTTTA	TCCCACGCCC AGGGTGCGGG
25851	TACATGTGGA ATGTACACCT	GTTACCAGCC CAATGGTCGG	ACAAATGGGA TGTTTACCCT	CTTGCGGCTG GAACGCCGAC	GAGCTGCCCA CTCGACGGGT
25901				CGCGGGACCC GCGCCCTGGG	CACATGATAT GTGTACTATA
25951	CCCGGGTCAA GGGCCCAGTT	CGGAATACGC GCCTTATGCG	GCCCACCGAA CGGGTGGCTT	ACCGAATTCT TGGCTTAAGA	CCTGGAACAG GGACCTTGTC
26001	GCGGCTATTA CGCCGATAAT	CCACCACACC GGTGGTGTGG	TCGTAATAAC AGCATTATTG	CTTAATCCCC GAATTAGGGG	GTAGTTGGCC CATCAACCGG
26051	CGCTGCCCTG GCGACGGGAC	GTGTACCAGG CACATGGTCC	AAAGTCCCGC TTTCAGGGCG	TCCCACCACT AGGGTGGTGA	GTGGTACTTC CACCATGAAG
26101	CCAGAGACGC GGTCTCTGCG	CCAGGCCGAA GGTCCGGCTT	GTTCAGATGA CAAGTCTACT	CTAACTCAGG GATTGAGTCC	GGCGCAGCTT CCGCGTCGAA
26151	GCGGGCGGC1	TTCGTCACAG AAGCAGTGTC	GGTGCGGTCC CCACGCCAGC	CCCGGGCAGG GGGCCCGTCC	GTATAACTCA CATATTGAGT
26201	CCTGACAATO GGACTGTTAO	AGAGGGCGAC TCTCCCGCTC	GTATTCAGCT CATAAGTCGA	CAACGACGAC GTTGCTGCTC	TCGGTGAGCT AGCCACTCGA
26251	CCTCGCTTG(GGAGCGAAC(TCTCCGTCCC	GACGGGACAT CTGCCCTGT	TTCAGATCG(A AAGTCTAGC(9009090090 90090909090
26301	CGCTCTTCAT GCGAGAAGTI	TCACGCCTC	G TCAGGCAATO C AGTCCGTTAO	CTAACTCTGG GGATTGAGACG	AGACCTCGTC TCTGGAGCAG
26351	CTCTGAGCC(GAGACTCGG	G CGCTCTGGA	G GCATTGGAA C CGTAACCTT	C TCTGCAATT G AGACGTTAA	T ATTGAGGAGT A TAACTCCTCA
26401	TTGTGCCAT	C GGTCTACTT G CCAGATGAA	T AACCCCTTC A TTGGGGAAG	T CGGGACCTC A GCCCTGGAG	C CGGCCACTAT G GCCGGTGATA
26451	CCGGATCAA GGCCTAGTT	T TTATTCCTA A AATAAGGAT	A CTTTGACGC T GAAACTGCG	G GTAAAGGAC C CATTTCCTG	T CGGCGGACGG A GCCGCCTGCC

Figure 27 AB

26501		ATGTTAAGTG TACAATTCAC			
26551		TCGCCGCCAC AGCGGCGGTG			
26601		AATTGCCCGA TTAACGGGCT			
26651		GCCCAGGGAG CGGGTCCCTC			
26701	GGGTCGCGGG	CCTGCTAGTT GGACGATCAA	CTCGCCCTGT	CCCCTGGGAC	ACAAGAGTGA
26751	CACTAAACGT	ACTGTCCTAA TGACAGGATT	GGGACCTAAT	GTAGTTCTAG	AAACAACGGT
26801	AGAGACACGA	GAGTATAATA CTCATATTAT	TTATGTCTTT	AATTTTATAT	GACCCCGAGG
26851	ATAGCGGTAG	CTGTAAACGC GACATTTGCG	GTGGCAGAAG	TGGGCGGGTT	CGTTTGGTTC
26901	CGCTTGGAAT	CCTGGTACTT GGACCATGAA	AATTGTAGAG	AGGGAGACAC	TAAATGTTGT
26951	CAAAGTTGGG	AGACGGAGTG TCTGCCTCAC	TCAGATGCTC	TCTTGGAGAG	GCTCGAGTCG
27001	ATGAGGTAGT	GAAAAAACAC CTTTTTTGTG	GTGGGAGGAA	TGGACGGCCC	TTGCATGCTC
27051	ACGCAGTGGC	GCCGCTGCAC	GTGTGGATGG	CGGACTGGCA	TTTGGTCTGA
27101	AAAAGGCCTG	TCTGGAGTTA	TTGAGACAAA	TGGTCTTGTC	GAGGTGAGCT CTCCACTCGA
	ATCTTTTGGG	AATCCCATAA	TCCGGTTTCC	GCGTCGATGA	GTGGGGTTTA CACCCCAAAT
	ACTTGTTAAG	TTCGTTGAGA	TGCCCGATAA	GATTAAGTCO	TTTCTCTAGA AAAGAGATCT
	TAGCCCCAAC	CCCAATAAGA	GACAGAACAC	TAAGAGAAAT	TTCTTATACT AAGAATATGA
	TTGCGAAGAC	ACGGATTCCG	AGCGGCGGAC	GACACACGTO	ATTTGCATTT TAAACGTAAA
	TAACAGTCGA	A AAAATTTGCG	ACCCCAGCGG	TGGGTTCTA	ATTAGGTACA TAATCCATGT
27401	TAATCCTAGO ATTAGGATCO	TTTACTCACC AAATGAGTGG	GAACGCAGTCAC	GGGTGCCAT(CACCCAAAAG GTGGGTTTTC

Figure 27AC

VO 02/0220	080				PCT/US01/28861
27451	GTGGATTT.	AGGAGCCAGC	CTGTAATGTT	ACATTCGC.	CTGAAGCTAA
		TCCTCGGTCG			- ·
27501	TGAGTGCACC	ACTCTTATAA	AATGCACCAC	AGAACATGAA	AAGCTGCTTA
	ACTCACGTGG	TGAGAATATT	TTACGTGGTG	TCTTGTACTT	TTCGACGAAT
27551	TTCGCCACAA	AAACAAAATT	GGCAAGTATG	CTGTTTATGC	TATTTGGCAG
	AAGCGGTGTT	TTTGTTTTAA	CCGTTCATAC	GACAAATACG	ATAAACCGTC
27601					•
		GATGTCTCAT			
27651	TAAAACTTTT				
		TACATATGAA			
27701	- -	CAAACAGTAT			
		GTTTGTCATA			
27751	AACACTGGCA				
07001					ACGAGCGAAA
27801	GGTCTGTACC	CTACTCTATA			
22051		AATGCCTTAA			
27851		TTACGGAATT			
27901		ACTCGCTGCT			
00000					AATCGTAATA
2/951	AATTAGAATA				TTATGGTAAG
2001					ACAACCTTGA
28001					TGTTGGAACT
28051					ACCTGTCCCG
				•	TGGACAGGGC
28101					GAGATGACCA CTCTACTGGT
28151					CACAAATACA
					GTGTTTATGT
28201					GCATGTGGTG
					CGTACACCAC
28251	GTTCTCCATA				
					ACCGAGTAGA
28301					TCCCATCATT
					AGGGTAGTAA
28351					GACTGAAACA
	CACGATGTGG	GTTTGTTACT	ACCTTAGGTA	TCTAACCTGC	CTGACTTTGT



28451		CTGACCCTTG GACTGGGAAC		
28501	CTGCGGTTTC GACGCCAAAG	TCACATCGAA AGTGTAGCTT		
28551		GATTTGTCAC CTAAACAGTG		-
28601		TTTATCCAGT AAATAGGTCA		
28651	ATCTCAGACA TAGAGTCTGT	CCATCCCCAG GGTAGGGGTC		
28701		AATTATGAAA TTAATACTTT		
28751		CGTTTTGTTC GCAAAACAAG		
28801		ACTCGTATAT TGAGCATATA		
28851		CGAAGCCTGG GCTTCGGACC		
28901		TCTTAGCCCT AGAATCGGGA		
28951		GATGCCATGA CTACGGTACT		
29001		ACAAGTTGTT TGTTCAACAA		
29051		CTCCCACCC GAGGGTGGGG		
29101				TATTACAGAG ATAATGTCTC
29151				AGCGCATGAA TCGCGTACTT
29201				AGGGGTATCT TCCCCATAGA
29251				TACCACCGGA ATGGTGGCCT
29301				TGGTGGTCAT ACCACCAGTA

Figure 27 A E

29401		CTCACCTTGT GAGTGGAACA		
29451	AAGACCCTGT TTCTGGGACA	GCGGTCTCAA CGCCAGAGTT		
29501	AATAATAAAG TTATTATTTC	CATCACTTAC GTAGTGAATG		
29551		GCACCTCCTT CGTGGAGGAA		
29601		GCAAACTTTC CGTTTGAAAG		
29651		TCCATCCGCA AGGTAGGCGT		
29701		CGTCTGAAGA GCAGACTTCT		
29751		CCTCCAACTG GGAGGTTGAC		
29801		TCAAGAGAGT AGTTCTCTCA		
29851		TTACCTCCAA AATGGAGGTT		
29901		GACGAGGCCG CTGCTCCGGC		
29951				GGAAATATCT CCTTTATAGA
30001	GCACCCCTCA CGTGGGGAGT			CCGCCGCACC
30051				GCCCCGCTAA CGGGGCGATT
30101				CCTCACAGTG GGAGTGTCAC
30151				CCACCACCGA GGTGGTGGCT
30201				ACTGCCACTG TGACGGTGAC
30251				AAATGGAAAA TTTACCTTTT

Figure 27 AF

30351		GCAACTGGTC CGTTGACCAG			
30401	AAACTAAAGT TTTGATTTCA	TACTGGAGCC ATGACCTCGG			
30451		CAGGAGGACT GTCCTCCTGA			
30501		AGTTATCCGT TCAATAGGCA		•	
30551		CCCTCTTTTT GGGAGAAAAA			
30601		GCCTTTACTT CGGAAATGAA			
30651		CTAAGCACTG GATTCGTGAC			
30701	ATCGGTAATT	TGCAGGAGAT ACGTCCTCTA	CCCGAACTTA	AACCAAGTGG	ATTACGTGGT
30751	TTGTGTTTAG	CCCTCAAAAC GGGAGTTTTG	TTTTTAACCG	GTACCGGATC	TTÄAACTAAG
30801	TTTGTTCCGA	ATGGTTCCTA TACCAAGGAT	TTGATCCTTG	ACCGGAATCA	AAACTGTCGT
30851		TACAGTAGGA ATGTCATCCT			
30901		CTCCATCTCC GAGGTAGAGG			
30951		TTGGTCTTAA AACCAGAATT			
31001					TGGAACAGTT ACCTTGTCAA
	GTTTCACGAG	TAGAATAATA	TTCTAAACTG	CTTTTACCTC	
31101		CTGGACCCAG GACCTGGGTC			GGAGATCTTA CCTCTAGAAT
31151		AGCCTATACA TCGGATATGT			TAACCTATCA ATTGGATAGT
31201					TTGTCAGTCA AACAGTCAGT

Figure 27 AG

31251		AACGGAGACA TTGCCTCTGT			ACCATTACAC TGGTAATGTG
31301		ACAGGAAACA TGTCCTTTGT			
31351		GGGACTGGTC CCCTGACCAG			
31401		TACACTTTTT ATGTGAAAAA			
31451		TCAACGTGTT AGTTGCACAA			
31501		AGTAGTATAG TCATCATATC			
31551		TCAAACTCAC AGTTTGAGTG			
31601	AGGGTTGTGT	CAGAGTACAC GTCTCATGTG	TCAGGAAAGA	GGGGCCGACC	GGAATTTTTC
31651	GTAGTATAGT	TGGGTAACAG ACCCATTGTC	TGTATAAGAA	TCCACAATAT	AAGGTGTGCC
31701	AAAGGACAGC		AGTAGTCACT	TTTAATTATT	GAGGGGCCCG
31751	TCGAGTGAAT	TCAAGTACAG	CGACAGGTCG	ACGACTCGGT	CAGGCTGCTG
31801	AGGTTGAACG	CCAACGAATT	GCCCGCCGCT	TCCTCTTCAG	CACGCCTACA GTGCGGATGT
31851	ACCCCCATCT	CAGTATTAGC	ACGTAGTCCT	ATCCCGCCAC	GTGCTGCAGC CACGACGTCG
	TCGCGCGCTT	ATTTGACGAC	GGCGGCGGCG	AGGCAGGACG	AGGAATACAA TCCTTATGTT
	GTACCGTCAC	CAGAGGAGTO	GCTACTAAGC	CTGGCGGGCG	AGCATAAGGC TCGTATTCCG
	CGGAACAGGA	GGCCCGTGTC	GTCGCGTGGG	ACTAGAGTGA	TAAATCAGCA ATTTAGTCGT
	GTCATTGACG	TCGTGTCGTG	GTGTTATAAC	: AAGTTTTAG(CACAGTGCAA G GTGTCACGTT
32101	GGCGCTGTAT CCGCGACATA	CCAAAGCTCA GGTTTCGAG1	TGGCGGGAC ACCGCCCTC	CACAGAACCO GTGTCTTGGO	ACGTGGCCAT G TGCACCGGTA
32151					T AAACACGCTG A TTTGTGCGAC

Figure 27 AH

32251		CTCTGATTAA GAGACTAATT		
32301		AACCTGCCCG TTGGACGGGC		
32351		AGTGGAGAGC TCACCTCTCG	•	
32401	••••	TCAATGTTGG AGTTACAACC	 	
32451		AAGCTCCTCC TTCGAGGAGG	 	
32501	-	TCAGCGTAAA AGTCGCATTT		
32551		TGCATTGTCA ACGTAACAGT		
32601		GGTAGCGCGG CCATCGCGCC		
32651	-	GAGTGCGCCG CTCACGCGGC		
32701		GGAACGCCGG CCTTGCGGCC		
32751	=	GACAAACAGA CTGTTTGTCT		
32801		TAGTTGTAGT ATCAACATCA		
32851		GGGTTCTATG CCCAAGATAC		
32901				CACATTCGTT GTGTAAGCAA
32951				ACCATGTTTT TGGTACAAAA
33001		CCAAAAGATT GGTTTTCTAA		GATCTATTAA CTAGATAATT
33051				GCCAAAGAAC CGGTTTCTTG
33101	AGATAATGGC TCTATTACCG			AAGGCAAACG TTCCGTTTGC

Figure 27 AI

33201	CTCTATAAAC GAGATATTTG				
33251	GCCACCTTCT CGGTGGAAGA	CAATATATCT GTTATATAGA			
33301	ATTGTAAAAA TAACATTTTT	TCTGCTCCAG AGACGAGGTC			
33351	AATCATGATT TTAGTACTAA	GCAAAAATTC CGTTTTTAAG			
33401		TTAACAAAAA AATTGTTTTT			
33451		ATAATCGTGC TATTAGCACG			
33501		CCATGACAAA GGTACTGTTT			
33551	CGGAGÇTATG GCCTCGATAC	CTAACCAGCG GATTGGTCGC	TAGCCCCGAT ATCGGGGCTA	GTAAGCTTGT CATTCGAACA	TGCATGGGCG ACGTACCCGC
33601		ATGCAAGGTG TACGTTCCAC			
33651		GCACATCGTA CGTGTAGCAT			
33701		ACCACAGAAA TGGTGTCTTT			AACATGTCTG TTGTACAGAC
33751	CGGGTTTCTG GCCCAAAGAC	CATAAACACA GTATTTGTGT	AAATAAAATA TATTTTATTT	ACAAAAAAAC TGTTTTTTG	ATTTAAACAT TAAATTTGTA
33801					CATAAGACGG GTATTCTGCC
33851	ACTACGGCCA TGATGCCGGT	TGCCGGCGTG ACGGCCGCAC	ACCGTAAAAA TGGCATTTTT	AACTGGTCAC TTGACCAGTG	CGTGATTAAA GCACTAATTT
33901	AAGCACCACC TTCGTGGTGG	GACAGCTCCT CTGTCGAGGA	CGGTCATGTC	CGGAGTCATA CCCTCAGTAT	ATGTAAGACT TACATTCTGA
33951	CGGTAAACAC GCCATTTGTG	ATCAGGTTGA TAGTCCAACT	TTCACATCGG	TCAGTGCTAA AGTCACGATT	AAAGCGACCG TTTCGCTGGC
34001	AAATAGCCCG TTTATCGGGC	GGGGAATACA CCCCTTATGT	TACCCGCAGG	GCATCTCTGT	ACATTACAGC TGTAATGTCG
34051	CCCCATAGGA GGGGTATCCI	GGTATAACAA CCATATTGTT	AATTAATAGO TTAATTATCO	AGAGAAAAAC TCTCTTTTTC	ACATAAACAC G TGTATTTGTG

Figure 27AJ

34151				ACAGTCAGCC TGTCAGTCGG	
34201	AAAAGAAAAC TTTTCTTTTG			GACACGGCAC CTGTGCCGTG	
34251				AGCGAGTATA TCGCTCATAT	
34301	AAAAATGACG TTTTTACTGC			AAACACCCAG TTTGTGGGTC	
34351	GCGAACCTAC CGCTTGGATG			AAACCCACAA TTTGGGTGTT	
34401				CTTCCCATTT GAAGGGTAAA	
34451				CCTAAAACCT GGATTTTGGA	
34501				AACTCCACCC TTGAGGTGGG	
			•		PacI
34551	ATATTGGCTT TATAACCGAA			TATTGATGAT	
	TATAACCGAA	GIIAGGIIII	ATTCCATATA	ATAACTACTA	CAMITAMITE
34601	AATTCGGATC	TGCGACGCGA	GGCTGGATGG	CCTTCCCCAT GGAAGGGGTA	TATGATTCTT
34601 34651	AATTCGGATC TTAAGCCTAG CTCGCTTCCG	TGCGACGCGA ACGCTGCGCT GCGGCATCGG	GGCTGGATGG CCGACCTACC GATGCCCGCG	CCTTCCCCAT	TATGATTCTT ATACTAAGAA TGCTGTCCAG
	AATTCGGATC TTAAGCCTAG CTCGCTTCCG GAGCGAAGGC GCAGGTAGAT	TGCGACGCGA ACGCTGCGCT GCGGCATCGG CGCCGTAGCC GACGACCATC	GGCTGGATGG CCGACCTACC GATGCCCGCG CTACGGGCGC	CCTTCCCCAT GGAAGGGGTA TTGCAGGCCA	TATGATTCTT ATACTAAGAA TGCTGTCCAG ACGACAGGTC CAAAAGGCCA
34651 34701	AATTCGGATC TTAAGCCTAG CTCGCTTCCG GAGCGAAGGC GCAGGTAGAT CGTCCATCTA	TGCGACGCGA ACGCTGCGCT GCGGCATCGG CGCCGTAGCC GACGACCATC CTGCTGGTAG AAAGGCCGCG	GGCTGGATGG CCGACCTACC GATGCCCGCG CTACGGGCGC AGGGACAGCT TCCCTGTCGA	CCTTCCCCAT GGAAGGGGTA TTGCAGGCCA AACGTCCGGT TCAAGGCCAG AGTTCCGGTC	TATGATTCTT ATACTAAGAA TGCTGTCCAG ACGACAGGTC CAAAAGGCCA GTTTTCCGGT GCTCCGCCCC
34651 34701 34751	AATTCGGATC TTAAGCCTAG CTCGCTTCCG GAGCGAAGGC GCAGGTAGAT CGTCCATCTA GGAACCGTAA CCTTGGCATT	TGCGACGCGA ACGCTGCGCT GCGGCATCGG CGCCGTAGCC GACGACCATC CTGCTGGTAG AAAGGCCGCG TTTCCGGCGC	GGCTGGATGG CCGACCTACC GATGCCCGCG CTACGGGCGC AGGGACAGCT TCCCTGTCGA TTGCTGGCGT AACGACCGCA	CCTTCCCCAT GGAAGGGGTA TTGCAGGCCA AACGTCCGGT TCAAGGCCAG AGTTCCGGTC TTTTCCATAG AAAAGGTATC	TATGATTCTT ATACTAAGAA TGCTGTCCAG ACGACAGGTC CAAAAGGCCA GTTTTCCGGT GCTCCGCCCC CGAGGCGGGG
34651 34701 34751 34801	AATTCGGATC TTAAGCCTAG CTCGCTTCCG GAGCGAAGGC GCAGGTAGAT CGTCCATCTA GGAACCGTAA CCTTGGCATT CCTGACGAGC GGACTGCTCG	TGCGACGCGA ACGCTGCGCT GCGGCATCGG CGCCGTAGCC GACGACCATC CTGCTGGTAG AAAGGCCGCG TTTCCGGCGC ATCACAAAAA TAGTGTTTTT	GGCTGGATGG CCGACCTACC GATGCCCGCG CTACGGGCGC AGGGACAGCT TCCCTGTCGA TTGCTGGCGT AACGACCGCA TCGACGCTCA AGCTGCGAGT	CCTTCCCCAT GGAAGGGGTA TTGCAGGCCA AACGTCCGGT TCAAGGCCAG AGTTCCGGTC TTTTCCATAG AAAAGGTATC AGTCAGAGGT TCAGTCTCCA	TATGATTCTT ATACTAAGAA TGCTGTCCAG ACGACAGGTC CAAAAGGCCA GTTTTCCGGT GCTCCGCCCC CGAGGCGGGG GGCGAAACCC CCGCTTTGGG TCCCTCGTGC
34651 34701 34751 34801 34851	AATTCGGATC TTAAGCCTAG CTCGCTTCCG GAGCGAAGGC GCAGGTAGAT CGTCCATCTA GGAACCGTAA CCTTGGCATT CCTGACGAGC GGACTGCTCG GACAGGACTA CTGTCCTGAT CTGTCCTGAT	TGCGACGCGA ACGCTGCGCT GCGGCATCGG CGCCGTAGCC GACGACCATC CTGCTGGTAG AAAGGCCGCG TTTCCGGCGC ATCACAAAA TAGTGTTTT TAAAGATACC ATTTCTÄTGG TCCGACCCTG	GGCTGGATGG CCGACCTACC GATGCCCGCG CTACGGGCGC AGGGACAGCT TCCCTGTCGA TTGCTGGCGT AACGACCGCA AGCTGCGAGT AGCTGCGAGT CGACGCTCA AGCTGCGAGT CCGCAAAGG CCGCTTACCG	CCTTCCCAT GGAAGGGTA TTGCAGGCCA AACGTCCGGT TCAAGGCCAG AGTTCCGTC TTTTCCATAG AAAAGGTATC AGTCAGAGGT TCAGTCTCCA CCCTGGAAGC GGGACCTTCG	TATGATTCTT ATACTAAGAA TGCTGTCCAG ACGACAGGTC CAAAAGGCCA GTTTTCCGGT GCTCCGCCCC CGAGGCGGGG GGCGAAACCC CCGCTTTGGG TCCCTCGTGC AGGGAGCACG CGCCTTTCTC
34651 34701 34751 34801 34851 34901	AATTCGGATC TTAAGCCTAG CTCGCTTCCG GAGCGAAGGC GCAGGTAGAT CGTCCATCTA GGAACCGTAA CCTTGGCATT CCTGACGAGC GGACTGCTCG GACAGGACTA CTGTCCTGAT CTGTCCTGAT CGAGAGGACA	TGCGACGCGA ACGCTGCGCT GCGGCATCGG CGCCGTAGCC GACGACCATC CTGCTGGTAG AAAGGCCGCG ATCACAAAAA TAGTGTTTT TAAAGATACC ATTTCTATGG TCCGACCCTG AGGCTGGGAC	GGCTGGATGG CCGACCTACC GATGCCCGCG CTACGGGCGC AGGGACAGCT TCCCTGTCGA TTGCTGGCGT AACGACCGCA AGCTGCGAGT AGCTGCGAGT CCGCAAAGG CCGCTTACCG GGCGAATGGC TTCTCATAGC	CCTTCCCCAT GGAAGGGGTA TTGCAGGCCA AACGTCCGGT TCAAGGCCAG AGTTCCGGTC TTTTCCATAG AAAAGGTATC AGTCAGAGGT TCAGTCTCCA CCCTGGAAGC GGGACCTTCG GATACCTGTC CTATGGACAG	TATGATTCTT ATACTAAGAA TGCTGTCCAG ACGACAGGTC CAAAAGGCCA GTTTTCCGGT GCTCCGCCCC CGAGGCGGGG GGCGAAACCC CCGCTTTGGG TCCCTCGTGC AGGGAGCACG CGCCTTTCTC GCGGAAAGAG GGTATCTCAG

Figure 27 AK

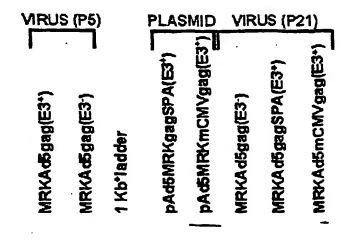
	AAGTCGGGCT	GGCGACGCGG	AATAGGCCAT	TGATAGCAGA	ACTCAGGTTG
35101	CCCCTAAGAC	ACCACTTATC	CCCACTCCCA	GCAGCCACTG	CTAACACCAT
33101					
	GGCCATTCTG	IGCIGAATAG	CGGTGACCGT	CGTCGGTGAC	CATTGTCCTA
35151	TAGCAGAGCG	AGGTATGTAG	GCGGTGCTAC	AGAGTTCTTG	AAGTGGTGGC
	ATCGTCTCGC	TCCATACATC	CGCCACGATG	TCTCAAGAAC	TTCACCACCG
35201	CTAACTACGG	CTACACTAGA	AGGACAGTAT	TTGGTATCTG	CGCTCTGCTG
				AACCATAGAC	
		wildlanici		in contrast	oc on one one
35251	AAGCCAGTTA	ССТТСССААА	AACACTTCCT	AGCTCTTGAT	CCCCCAAACA
JJ2J2				TCGAGAACTA	
	110010,011	odmocc111	11C1CIMCCI	1 CONGRETA	0000011101
35301	AACCACCGCT	CCTACCCCTC	CUTTUTTUTT	TTGCAAGCAG	САСАТТАССС
33301				AACGTTCGTC	
	1100100CGA	CCATCOCCAC	CARAMARICA	MCGIICGIC	GICIANIGCO
35351	CCDCDDDDDD	ACCATOTOAA	СВВСВТССТТ	TGATCTTTTC	тассссстст
33331	••			ACTAGAAAAG	
	CGICIIIIII	ICCINGAGII	CIICIAGGAA	ACIAGAAAAG	AIGCCCCAGA
35401	CACCCTCACT	CCAACCAAAA	СТСАССТТАА	GGGATTTTGG	ጥር አጥር አር አጥጥ
33401			•	CCCTAAAACC	
	CIGCGAGICA	CCIIGCIIII	GNGIGCAAII	CCCIAAAACC	AGIACICIAA
35451	30C33333CC	አጥርጥጥር አርርጥ	እ C እ ጥርርጥጥጥ	AAATCAATCT	<u>አአአርጥአጥአጥአ</u>
33431				TTTAGTTAGA	
	IAGITITICC	INGANGIGGN	ICIAGGAAAA	IIINGIINGA	IIICAIAIAI
35501	ጥ ር እርጥ እ እ እርጥ	тестстелел	CTTACCAATC	CTTAATCAGT	GAGGCACCTA
33301				GAATTAGTCA	
	ACICATITOA	ACCAGACIGI	CAMIGGIIAC	GANTINGICA	CICCOIOGAI
35551	יירייר א כרכי איי	C 中C中C中 2 中中中	CCTTCATCCA	TAGTTGCCTG	ACTCCCCCTC
33331				ATCAACGGAC	
	AGAGICGCIA	GUCUGUIVA	GCAAGTAGGT	ATCAACGGAC	randadacno
35601	GTGTAGATAA	CTACGATACG	GGAGGGCTTA	CCATCTGGCC	CCAGTGCTGC
JJ001				GGTAGACCGG	
	Chemicinii		••••••••		
35651	AATGATACCG	CGAGACCCAC	GCTCACCGGC	TCCAGATTTA	TCAGCAATAA
				AGGTCTAAAT	
					•
35701	ACCAGCCAGC	CGGAAGGGCC	GAGCGCAGAA	GTGGTCCTGC	AACTTTATCC
••••				CACCAGGACG	
•					
35751	GCCTCCATCC	AGTCTATTAA	TTGTTGCCGG	GAAGCTAGAG	TAAGTAGTTC
				CTTCGATCTC	
35801	GCCAGTTAAT	AGTTTGCGCA	ACGTTGTTGC	CATTGCTACA	GGCATCGTGG
				GTAACGATGT	
		••••			
35851	TGTCACGCTC	GTCGTTTGGT	ATGGCTTCAT	TCAGCTCCGG	TTCCCAACGA
				AGTCGAGGCC	
35901	TCAAGGCGAG	TTACATGATC	CCCCATGTTG	TGCAAAAAAG	CGGTTAGCTC
				ACGTTTTTTC	
35951	CTTCGGTCCT	CCGATCGTTG	TCAGAAGTAA	GTTGGCCGCA	GTGTTATCAC
				CAACCGGCGT	
				2.2.2.2.444.	

Figure 2 7AL

36051	AGATGCTTTT	CTGTGACTGG	TGAGTACTCA	ACCAAGTCAT	TCTGAGAATA
	TCTACGAAAA	GACACTGACC	ACTCATGAGT	TGGTTCAGTA	AGACTCTTAT
36101	GTGTATGCGG	CGACCGAGTT	GCTCTTGCCC	GGCGTCAACA	CGGGATAATA
	CACATACGCC	GCTGGCTCAA	CGAGAACGGG	CCGCAGTTGT	GCCCTATTAT
36151	CCGCGCCACA	TAGCAGAACT	TTAAAAGTGC	TCATCATTGG	AAAACGTTCT
	GGCGCGGTGT	ATCGTCTTGA	AATTTTCACG	AGTAGTAACC	TTTTGCAAGA
36201	TCGGGGCGAA	AACTCTCAAG	GATCTTACCG	CTGTTGAGAT	CCAGTTCGAT
	AGCCCCGCTT	TTGAGAGTTC	CTAGAATGGC	GACAACTCTA	GGTCAAGCTA
36251				AGCATCTTTT	
	CATTGGGTGA	GCACGTGGGT	TGACTAGAAG	TCGTAGAAAA	TGAAAGTGGT
36301				AAAATGCCGC	
	•			TTTTACGGCG	
36351				ATACTCTTCC	
				TATGAGAAGG	
36401				CATGAGCGGA	
				GTACTCGCCT	
36451				TTCCGCGCAC	
				AAGGCGCGTG	
36501				ATTATCATGA	
				TAATAGTACT	
36551				TCTTCAAGAA	
	ATTTTTATCC	GCATAGTGCT	CCGGGAAAGC	AGAAGTTCTT	AACCTAGGCT
		PacI			
		~~~~~~			

36601 ATTCTTAATT TCTTAATTAA (SEQ ID NO:34)
TAAGAATTAA AGAATTAATT (SEQ ID NO:35)

Ingure 27 AM



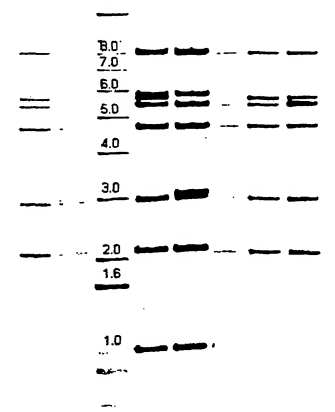


FIGURE 28

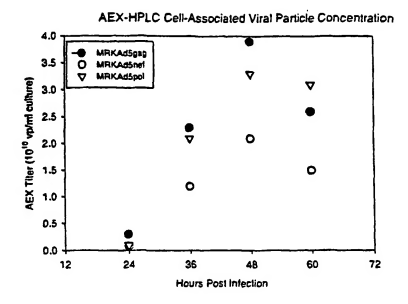


FIGURE 29A

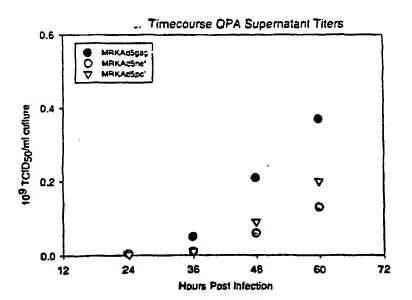


FIGURE 29B

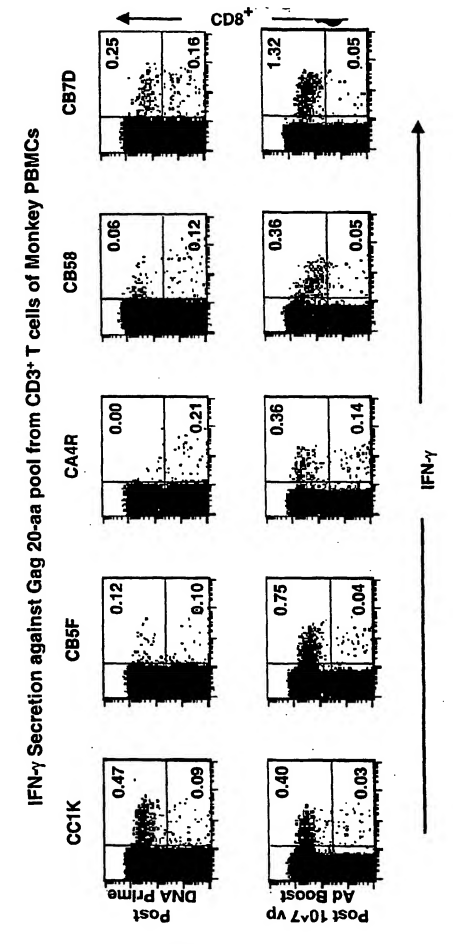
atg Met 1	gat Asp	gca Ala	atg Met	aag Lys 5	aga Arg	GJA aaa	ctc Leu	tgc Cys	tgt Cys 10	gtg Val	ctg Leu	ctg Leu	ctg Leu	tgt Cys 15	Gja āāa	48
gca Ala	gtc Val	ttc Phe	gtt Val 20	tcg Ser	ccc Pro	agc Ser	gag Glu	atc Ile 25	tcc Ser	att Ile	gtg Val	tgg Trp	gcc Ala 30	tcc Ser	agg Arg	96
gag Glu	ctg Leu	gag Glu 35	agg Arg	ttt Phe	gct Ala	gtg Val	aac Asn 40	cct Pro	ggc Gly	ctg Leu	ctg Leu	gag Glu 45	acc Thr	tct Ser	gag Glu	144
ggg Gly	tgc Cys 50	agg Arg	cag Gln	atc Ile	ctg Leu	ggc Gly 55	cag Gln	ctc Leu	cag Gln	ccc Pro	tcc Ser 60	ctg Leu	caa Gln	aca Thr	Gly	192
tct Ser 65	gag Glu	gag Glu	ctg Leu	agg Arg	tcc Ser 70	ctg Leu	tac Tyr	aac Asn	aca Thr	gtg Val 75	gct Ala	acc Thr	ctg Leu	tac Tyr	tgt Cys 80	240
gtg Val	cac His	cag Gln	aag Lys	att Ile 85	gat Asp	gtg Val	aag Lys	gac Asp	acc Thr 90	aag Lys	gag Glu	gcc Ala	ctg Leu	gag Glu 95	aag Lys	288
att Ile	gag Glu	gag Glu	gag Glu 100	cag Gln	aac Asn	aag Lys	tcc Ser	aag Lys 105	Lys	aag Lys	gcc Ala	cag Gln	cag Gln 110	gct Ala	gct Ala	336
gct Ala	ggc Gly	aca Thr 115	ggc	aac Asn	tcc Ser	agc Ser	cag Gln 120	Val	tcc Ser	cag Gln	aac Asn	tac Tyr 125	Pro	att	gtg Val	384
cag Gln	aac Asn 130	Leu	cag Gln	ggc	cag Gln	atg Met 135	Val	cac His	cag Gln	gcc	ato Ile 140	Ser	Pro	cgg Arg	acc	432
ctg Lev 145	a Asn	gcc Ala	tgg Trp	gtg Val	aag Lys 150	Val	gtg Val	gag Glu	g gag 1 Glu	aag Lys 155	Ala	tto Phe	tco Ser	e cct	gag Glu 160	480
gto Val	ato l Ile	ccc Pro	atg Met	ttc Phe 165	Ser	gcc	ctg Lev	tct Ser	gag Glv 170	ı Gly	gco Ala	e aco	c ccc	Cag Glr 175	Asp	528
Ct: Lei	g aac ı Asr	aco Thi	atg Met	Let	aac Asn	aca Thr	gtg Val	999 1 Gly 18!	λ C17	cat / His	caq Gli	g gc n Ala	gco Ala 19	Me	g cag	576
at: Me	g cto t Lei	aaq 1 Ly: 19:	s Gli	aco Thi	ato Ile	aat Asi	gag 1 Glu 200	ı Gl	g gci	t gct a Ala	gag a Gl	g tg u Tr 20	p As	age p Are	g ctg g Leu	624
ca Hi	t cci s Pro 210	va.	g cad	gct Ala	ggc Gly	21!	o Ile	t gc	c cc	c ggo o Gl	c ca y Gl 22	n Me	g ag t Ar	g ga g Gl	g ccc	672
ag Ar 22	g Gl	c tc y Se:	t gad r Asj	e att	e Ala 23	a G1;	c ac	c ac r Th	c tc r Se	c acc r Thi	r Le	c ca u Gl	g ga n Gl	g ca u Gl	g att n Ile 240	720
99 G1	c tg y Tr	g at p Me	g aco	c aa r As: 24	n Ası	c cc n Pr	c cc o Pr	c at o Il	c cc e Pr 25	o va	g gg 1 Gl	y Gl	a at u Il	c ta e Ty 25	c aag r Lys 5	768

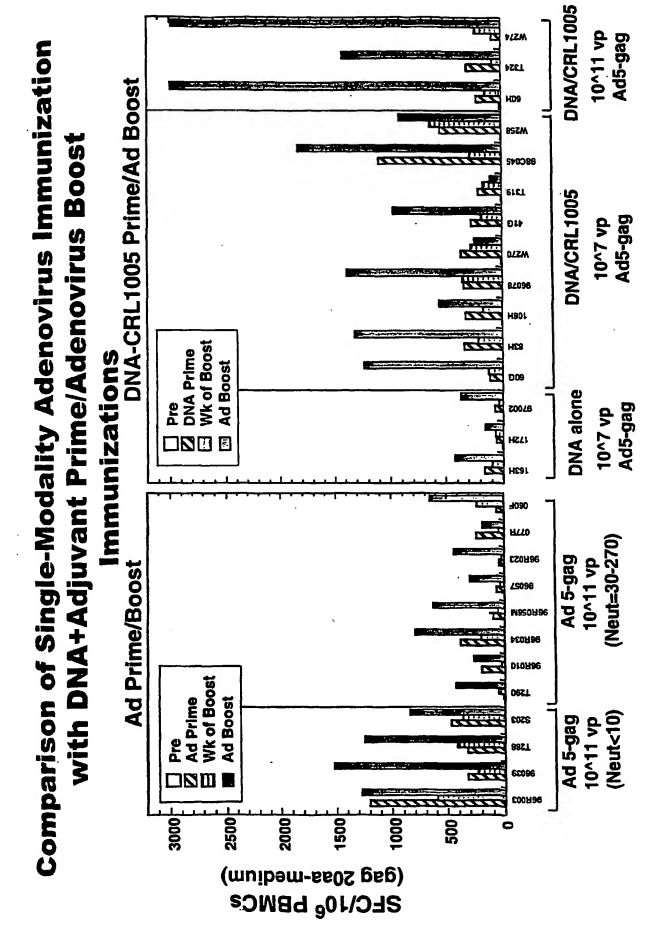
Figure 30'A"

													tac Tyr 270			816
													ttc Phe			864
													gcc Ala			912
gag Glu 305	gtg Val	aag Lys	aac Asn	jib	atg Met 310	aca Thr	gag Glu	acc Thr	ctg Leu	ctg Leu 315	gtg Val	cag Gln	aat Asn	gcc Ala	aac Asn 320	960
													gcc Ala			1008
													ggt Gly 350			1056
gcc Ala	agg Arg	gtg Val 355	ctg Leu	gct Ala	gag Glu	gcc Ala	atg Met 360	tcc Ser	cag Gln	gtg Val	acc Thr	aac Asn 365	tcc Ser	gcc Ala	acc Thr	1104
												Lys	aca Thr			1152
											Lys		tgt Cys			1200
										Lys			cac His			1248
rys Pag	gac Asp	tgc Cys	aat Asn 420	gag Glu	agg Arg	cag Gln	gcc Ala	aac Asn 425	ttc Phe	ctg Leu	Gly	aaa Lys	atc Ile 430	Trp	Pro	1296
								Phe					cct Pro			1344
aca Thr	gcc Ala 450	Pro	Pro	gag Glu	gag Glu	tcc Ser 455	Phe	agg Arg	ttt Phe	6JÀ	gag Glu 460	Glu	aag Lys	acc	Thr	1392
	Ser					Pro					Lev		Pro			1440
tcc Ser	ctg Leu	agg Arg	tcc Ser	ctg Leu 485	Phe	Gly	aac Asn	gac	Pro 490	Ser	tcc Ser	Glr.	taa	(SI (SI	D NO:36) D NO:37)	1482



Figure 31





## FIGURE 33A

አጥርርርጥርርጥል	GGGCTTCTGT	<b>CCTCTCTCCT</b>	GGTGAGCTGG	ACAAGTGGGA	GAAGATCAGG
	GTGGCAAGAA				
	TTGCTGTGAA		•		
	TCCAGCCCTC				
	CCCTGTACTG				
	TTGAGGAGGA				
	ACTCCAGCCA				
	ACCAGGCCAT				
	TCTCCCCTGA				
	TGAACACCAT				
	CCATCAATGA				
	CCCCCGGCCA				
	AGGAGCAGAT				
	GGTGGATCAT				
	ACATCAGGCA				
	TGAGGGCTGA				
	AGAATGCCAA				
	AGGAGATGAT				
	CTGAGGCCAT				
	GGAACCAGAG				
	ACTGTAGGGC				
	ACTGTAGGGC				
	GGCCTGGCAA				
	GGCCTGGCAA GGTTTGGGGA				
	ACCCCCTGGC				
	TCTCCCCCAT				
	AGCAGTGGCC				
					CTACAACACC
					GGACTTCAGG
					CCACCCCGCT
					CTTCTCTGTG
					CAACAATGAG
					CTCCCCTGCC
					CCCTGACATT
					TGGGCAGCAC
					CACCCCTGAC
					A CCCCGACAAG
					A TGACATCCAG
					A GGTGAGGCAG
					CACTGAGGAG
					A TGGGGTGTAC
GC TGAGCTG	S MGCIGGCIG	ומטטטאטאט ב	S MICCIGANG	- NGCCIGIGG	

### FIGURE 33B

TATGACCCCT	CCAAGGACCT	GATTGCTGAG	ATCCAGAAGC	AGGGCCAGGG	CCAGTGGACC
TACCAAATCT	ACCAGGAGCC	CTTCAAGAAC	CTGAAGACTG	GCAAGTATGC	CAGGATGAGG
GGGGCCCACA	CCAATGATGT	GAAGCAGCTG	ACTGAGGCTG	TGCAGAAGAT	CACCACTGAG
TCCATTGTGA	TCTGGGGCAA	GACCCCCAAG	TTCAAGCTGC	CCATCCAGAA	GGAGACCTGG
GAGACCTGGT	GGACTGAGTA	CTGGCAGGCC	ACCTGGATCC	CTGAGTGGGA	GTTTGTGAAC
ACCCCCCCC	TGGTGAAGCT	GTGGTACCAG.	CTGGAGAAGG	AGCCCATTGT	GGGGGCTGAG
ACCTTCTATG	TGGCTGGGGC	TGCCAACAGG	GAGACCAAGC	TGGGCAAGGC	TGGCTATGTG
ACCAACAGGG	GCAGGCAGAA	GGTGGTGACC	CTGACTGACA	CCACCAACCA	GAAGACTGCC
CTCCAGGCCA	TCTACCTGGC	CCTCCAGGAC	TCTGGCCTGG	AGGTGAACAT	TGTGACTGCC
TCCCAGTATG	CCCTGGGCAT	CATCCAGGCC	CAGCCTGATC	AGTCTGAGTC	TGAGCTGGTG
AACCAGATCA	TTGAGCAGCT	GATCAAGAAG	GAGAAGGTGT	ACCTGGCCTG	GGTGCCTGCC
CACAAGGGCA	TTGGGGGCAA	TGAGCAGGTG	GACAAGCTGG	TGTCTGCTGG	CATCAGGAAG
GTGCTGTTCC	TGGATGGCAT	TGACAAGGCC	CAGGATGAGC	ATGAGAAGTA	CCACTCCAAC
TGGAGGGCTA	TGGCCTCTGA	CTTCAACCTG	CCCCCTGTGG	TGGCTAAGGA	GATTGTGGCC
TCCTGTGACA	AGTGCCAGCT	GAAGGGGGAG	GCCATGCATG	GGCAGGTGGA	CTGCTCCCCT
GGCATCTGGC	AGCTGGCCTG	CACCCACCTG	GAGGGCAAGG	TGATCCTGGT	GGCTGTGCAT
GTGGCCTCCG	GCTACATTGA	GGCTGAGGTG	ATCCCTGCTG	AGACAGGCCA	GGAGACTGCC
TACTTCCTGC	TGAAGCTGGC	TGGCAGGTGG	CCTGTGAAGA	CCATCCACAC	TGCCAATGGC
TCCAACTTCA	CTGGGGCCAC	AGTGAGGGCT	GCCTGCTGGT	GGGCTGGCAT	CAAGCAGGAG
TTTGGCATCC	CCTACAACCC	CCAGTCCCAG	GGGGTGGTGG	CCTCCATGAA	CAAGGAGCTG
AAGAAGATCA	TTGGGCAGGT	GAGGGACCAG	GCTGAGCACC	TGAAGACAGO	TGTGCAGATG
GCTGTGTTCA	TCCACAACTT	CAAGAGGAAG	GGGGGCATCG	GGGGCTACTC	CGCTGGGGAG
AGGATTGTGG	ACATCATTGC	CACAGACATC	CAGACCAAGG	AGCTCCAGAA	GCAGATCACC
AAGATCCAGA	ACTTCAGGGT	GTACTACAGG	GACTCCAGGA	ACCCCTGTG	GAAGGGCCCT
-					TGACATCAAG
GTGGTGCCCA	GGAGGAAGGC	CAAGATCATC	AGGGACTATG	GCAAGCAGAT	GGCTGGGGAT
GACTGTGTGG	CCTCCAGGCA	GGATGAGGAC	TAA		
SEQ ID NO:	38				

#### FIGURE 34A

Met Gly Ala Arg Ala Ser Val Leu Ser Gly Gly Glu Leu Asp Lys Trp Glu Lys Ile Arg Leu Arg Pro Gly Gly Lys Lys Lys Tyr Lys Leu Lys His Ile Val Trp Ala Ser Arg Glu Leu Glu Arg Phe Ala Val Asn Pro Gly Leu Leu Glu Thr Ser Glu Gly Cys Arg Gln Ile Leu Gly Gln Leu Gln Pro Ser Leu Gln Thr Gly Ser Glu Glu Leu Arg Ser Leu Tyr Asn Thr Val Ala Thr Leu Tyr Cys Val His Gln Lys Ile Asp Val Lys Asp Thr Lys Glu Ala Leu Glu Lys Ile Glu Glu Glu Gln Asn Lys Ser Lys Lys Lys Ala Gln Gln Ala Ala Ala Gly Thr Gly Asn Ser Ser Gln Val Ser Gln Asn Tyr Pro Ile Val Gln Asn Leu Gln Gly Gln Met Val His Gln Ala Ile Ser Pro Arg Thr Leu Asn Ala Trp Val Lys Val Val Glu Glu Lys Ala Phe Ser Pro Glu Val Ile Pro Met Phe Ser Ala Leu Ser Glu Gly Ala Thr Pro Gln Asp Leu Asn Thr Met Leu Asn Thr Val Gly Gly His Gln Ala Ala Met Gln Met Leu Lys Glu Thr Ile Asn Glu Glu Ala Ala Glu Trp Asp Arg Leu His Pro Val His Ala Gly Pro Ile Ala Pro Gly Gln Met Arg Glu Pro Arg Gly Ser Asp Ile Ala Gly Thr Thr Ser Thr Leu Gln Glu Gln Ile Gly Trp Met Thr Asn Asn Pro Pro Ile Pro Val Gly Glu Ile Tyr Lys Arg Trp Ile Ile Leu Gly Leu Asn Lys Ile Val Arg Met Tyr Ser Pro Thr Ser Ile Leu Asp Ile Arg Gln Gly Pro Lys Glu Pro Phe Arg Asp Tyr Val Asp Arg Phe Tyr Lys Thr Leu Arg Ala Glu Gln Ala Ser Gln Glu Val Lys Asn Trp Met Thr Glu Thr Leu Leu Val Gln Asn Ala Asn Pro Asp Cys Lys Thr Ile Leu Lys Ala Leu Gly Pro Ala Ala Thr Leu Glu Glu Met Met Thr Ala Cys Gln Gly Val Gly Gly Pro Gly His Lys Ala Arg Val Leu Ala Glu Ala Met Ser Gln Val Thr Asn Ser Ala Thr Ile Met Met Gln Arg Gly Asn Phe Arg Asn Gln Arg Lys Thr Val Lys Cys Phe Asn Cys Gly Lys Val Gly His Ile Ala Lys Asn Cys Arg Ala Pro Arg Lys Lys Gly Cys Trp Lys Cys Gly Lys Glu Gly His Gln Met Lys Asp Cys Asn Glu Arg Gln Ala Asn Phe Leu Gly Lys Ile Trp Pro Ser His Lys Gly Arg Pro Gly Asn Phe Leu Gln Ser Arg Pro Glu Pro Thr Ala Pro Pro Glu Glu Ser Phe Arg Phe Gly Glu Glu Lys Thr Thr Pro Ser Gln Lys Gln Glu Pro Ile Asp Lys Glu Leu Tyr Pro Leu Ala Ser Leu Arg Ser Leu Phe Gly Asn Asp Pro Ser Ser Gln Met Ala Pro Ile Ser Pro Ile Glu Thr Val Pro Val Lys Leu Lys Pro Gly Met Asp Gly Pro Lys Val Lys Gln Trp Pro Leu Thr Glu Glu Lys Ile Lys Ala Leu Val Glu Ile Cys Thr Glu Met Glu Lys Glu Gly Lys Ile Ser Lys Ile Gly Pro Glu Asn Pro Tyr Asn Thr Pro Val Phe Ala Ile Lys Lys Asp Ser Thr Lys Trp Arg Lys Leu Val Asp Phe Arg Glu Leu Asn Lys Arg Thr Gln Asp Phe Trp Glu Val Gln Leu Gly Ile Pro His Pro Ala Gly Leu Lys Lys Lys Ser Val Thr Val Leu Ala Val Gly Asp Ala Tyr Phe Ser Val Pro Leu Asp Glu Asp Phe Arg Lys Tyr Thr Ala Phe Thr Ile Pro Ser Ile Asn Asn Glu Thr Pro Gly Ile Arg Tyr Gln Tyr Asn Val Leu Pro Gln Gly Trp Lys Gly Ser Pro Ala Ile Phe Gln Ser Ser Met Thr Lys Ile Leu Glu Pro Phe Arg Lys Gln Asn Pro Asp Ile Val Ile Tyr Gln Tyr Met Ala Ala Leu Tyr Val Gly Ser Asp Leu Glu Ile Gly Gln His Arg Thr Lys Ile Glu Glu Leu Arg Gln His Leu Leu Arg Trp Gly Leu Thr Thr Pro Asp Lys Lys His Gln Lys Glu Pro Pro Phe Leu Trp Met Gly Tyr Glu Leu His Pro

#### FIGURE 34B

Asp Lys Trp Thr Val Gln Pro Ile Val Leu Pro Glu Lys Asp Ser Trp Thr Val Asn Asp Ile Gln Lys Leu Val Gly Lys Leu Asn Trp Ala Ser Gln Ile Tyr Pro Gly Ile Lys Val Arg Gln Leu Cys Lys Leu Leu Arg Gly Thr Lys Ala Leu Thr Glu Val Ile Pro Leu Thr Glu Glu Ala Glu Leu Glu Leu Ala Glu Asn Arg Glu Ile Leu Lys Glu Pro Val His Gly Val Tyr Tyr Asp Pro Ser Lys Asp Leu Ile Ala Glu Ile Gln Lys Gln Gly Gln Gly Gln Trp Thr Tyr Gln Ile Tyr Gln Glu Pro Phe Lys Asn Leu Lys Thr Gly Lys Tyr Ala Arg Met Arg Gly Ala His Thr Asn Asp Val Lys Gln Leu Thr Glu Ala Val Gln Lys Ile Thr Thr Glu Ser Ile Val Ile Trp Gly Lys Thr Pro Lys Phe Lys Leu Pro Ile Gln Lys Glu Thr Trp Glu Thr Trp Trp Thr Glu Tyr Trp Gln Ala Thr Trp Ile Pro Glu Trp Glu Phe Val Asn Thr Pro Pro Leu Val Lys Leu Trp Tyr Gln Leu Glu Lys Glu Pro Ile Val Gly Ala Glu Thr Phe Tyr Val Ala Gly Ala Ala Asn Arg Glu Thr Lys Leu Gly Lys Ala Gly Tyr Val Thr Asn Arg Gly Arg Gln Lys Val Val Thr Leu Thr Asp Thr Thr Asn Gln Lys Thr Ala Leu Gln Ala Ile Tyr Leu Ala Leu Gln Asp Ser Gly Leu Glu Val Asn Ile Val Thr Ala Ser Gln Tyr Ala Leu Gly Ile Ile Gln Ala Gln Pro Asp Gln Ser Glu Ser Glu Leu Val Asn Gln Ile Ile Glu Gln Leu Ile Lys Lys Glu Lys Val Tyr Leu Ala Trp Val Pro Ala His Lys Gly Ile Gly Gly Asn Glu Gln Val Asp Lys Leu Val Ser Ala Gly Ile Arg Lys Val Leu Phe Leu Asp Gly Ile Asp Lys Ala Gln Asp Glu His Glu Lys Tyr His Ser Asn Trp Arg Ala Met Ala Ser Asp Phe Asn Leu Pro Pro Val Val Ala Lys Glu Ile Val Ala Ser Cys Asp Lys Cys Gln Leu Lys Gly Glu Ala Met His Gly Gln Val Asp Cys Ser Pro Gly Ile Trp Gln Leu Ala Cys Thr His Leu Glu Gly Lys Val Ile Leu Val Ala Val His Val Ala Ser Gly Tyr Ile Glu Ala Glu Val Ile Pro Ala Glu Thr Gly Gln Glu Thr Ala Tyr Phe Leu Leu Lys Leu Ala Gly Arg Trp Pro Val Lys Thr Ile His Thr Ala Asn Gly Ser Asn Phe Thr Gly Ala Thr Val Arg Ala Ala Cys Trp Trp Ala Gly Ile Lys Gln Glu Phe Gly Ile Pro Tyr Asn Pro Gln Ser Gln Gly Val Val Ala Ser Met Asn Lys Glu Leu Lys Lys Ile Ile Gly Gln Val Arg Asp Gln Ala Glu His Leu Lys Thr Ala Val Gln Met Ala Val Phe Ile His Asn Phe Lys Arg Lys Gly Gly Jle Gly Gly Tyr Ser Ala Gly Glu Arg Ile Val Asp Ile Ile Ala Thr Asp Ile Gln Thr Lys Glu Leu Gln Lys Gln Ile Thr Lys Ile Gln Asn Phe Arg Val Tyr Tyr Arg Asp Ser Arg Asn Pro Leu Trp Lys Gly Pro Ala Lys Leu Leu Trp Lys Gly Glu Gly Ala Val Val Ile Gln Asp Asn Ser Asp Ile Lys Val Val Pro Arg Arg Lys Ala Lys Ile Ile Arg Asp Tyr Gly Lys Gln Met Ala Gly Asp Asp Cys Val Ala Ser Arg Gln Asp Glu Asp SEQ ID NO: 39

International application No.

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		1.00.000	
A. CLAS IPC(7) US CL	SIFICATION OF SUBJECT MATTER : C12N 15/86 : 435/456		
	International Patent Classification (IPC) or to both na	tional classification and IPC	
B. FIEL	DS SEARCHED		
	rumentation searched (classification system followed to 24/205.1, 207.1, 227.1, 233.1; 435/69.1, 69.3, 173.3		
Documentation	on searched other than minimum documentation to the	extent that such documents are included	l in the fields searched
	ta base consulted during the international search (namontimuation Sheet	e of data base and, where practicable, so	earch terms used)
C. DOCT	JMENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.
X 	WO 96/39178 (ERTL et al.) 12 December 1996 (12. and claims 1 and 5.	12.1996), see page 5, 6,10, 12, 13	1-3, 8-11, 18
Y	and claims 1 and 3.		4, 5, 13-17, 29-32, 34, 35, 37
x	US 6,019,978 A (ERTL et al.) 1 February 2000,(01/	/02/2000), see columns 2, 7 and 8.	1-3, 8-11, 18
Y			4, 5, 13-17, 29-32, 34, 35, 37
X,P	US 6,287,571 <b>B</b> (ERTL et al.) 11 September 200 and claim 1.	1 (11/09/2001), see columns 2, 7, 8	1, 9, 18
x	US 5,643,579A (HUNG et al.) 1 July 1997 (01/07/1	1997), see examples 1, 2, 25 and 26.	1-3, 8, 9-11, 18
Y			4,5,13-17, 29-32, 34, 35, 37
Y	WANG et al. The use of an E1-deleted, replication expressing the rabies virus glycoprotein for early va Journal of Virology (March 1997) Vol. 71, No. 5, p	ecination of mice against rabies virus.	1-3, 9-11, 13-18
Further	documents are listed in the continuation of Box C.	See patent family annex.	
•	pecial categories of cited documents:	"T" later document published after the in	nternational filing date or
"A" document defining the general state of the art which is not considered to be of particular relevance		priority date and not in conflict with understand the principle or theory u	nderlying the invention
"E" earlier application or patent published on or after the international filing considered novel or cannot be considered to involve an date step when the document is taken alone			dered to involve an inventive
"L" document which may throw doubts on priority claim(s) or which is cited "Y" document of particular relevance; the claimed invention can to establish the publication date of another citation or other special reason considered to involve an inventive step when the document (as specified) combined with one or more other such documents, such combination being obvious to a person skilled in the art			step when the document is such documents, such
	at referring to an oral disclosure, use, exhibition or other means at published prior to the international filing date but later than the	*&* document member of the same pater	nt family
	date claimed actual completion of the international search	Date of mailing of the international se	arch report
	2002 (06.02.2002)	19 HUU ZUUZ	$\int$
Ca	nailing address of the ISA/US minissioner of Patents and Trademarks	Authorized officer	alblins for
	x PCT shington, D.C. 20231	Ulrike Winkler, Ph.D.	V
1	o. (703)305-3230	Telephone No. 703-308-0196	1/1

Form PCT/ISA/210 (second sheet) (July 1998)

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ategory *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Υ	NATUK et al. Immunogenicity of recombinant human adenovirus -human immunodeficiency virus vaccines in chimpanzees. Aids Research and Human Retroviruses (1993) Vol. 9, No. 5, pp395-404, see material and methods.	1, 9, 29-32
Y	PREVEC et al. Immune response to HIV-1 gag antigens induced by recombinant adenovirus vectors in mice and rhesus macaque monkeys. Journal of Acquired Immune Deficincy Syndrome. (1991) Vol. 4, No. 6 pp. 568-76, see abstract.	1, 9, 29-32
Y	LORI et al. Rapid protection against human immunodeficiency virus type 1 (HIV-1) replication mediated by high efficiency non-retroviral delivery of genes interfering with HIV-1 tat and gag. Gene Therapy (1994) Vol. 1, No. 1, pp. 27-31, see abstract.	1, 9
Y	PFARR et al. Differential effects of polyadenylation regions on gene expression in mammalian cells. DNA (1986) Vol. 5, No. 2, pp.115-22, see abstract.	16
Y	NATUK et al. Adenovirus vectored vaccine. Developmental Biological Standards (1994) Vol. 82, pp. 71-77, see abstract.	1, 9

International application No.

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Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)					
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:					
1.		Claim Nos.: because they relate to subject matter not required to be searched by this Authority, namely:			
2.		Claim Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:			
3.	6.4(a).	Claim Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule			
Box	II Ob	servations where unity of invention is lacking (Continuation of Item 2 of first sheet)			
		tional Searching Authority found multiple inventions in this international application, as follows:			
1.		As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.			
2.		As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.			
3.		As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:			
4.	$\boxtimes$	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-5, 8-11, 13-18, 29-32, 34, 35, 37			
Ren	nark on	Protest			

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## BOX II. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING

This application contains the following inventions or groups of inventions which are not so linked as to form a single general inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.

Group	Claims	
1	1-5, 8-11, 13-18, 29, 30, 31, 32, 34, 35, 37	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Gag protein (SEQ ID NO: 29) inserted in the parallel orientation of E1. In addition the vector contains a promoter and a polyadenylation signal.
2	6, 7, 36	The claims are directed to an adenoviral vector that is at least partially deleted of <u>AE1</u> and <u>AE3</u> , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Gag protein (SEQ ID NO: 29).
3	12, 33	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV protein inserted in the antiparallel orientation of E1.
4	19-23, 38-42	The claims are directed to a method of making and harvesting of a recombinant adenoviral particle that contains a gene encoding an HIV Gag protein.
5	24, 27, 28, 43, 46, 47	The claim is directed to a method of generating a cellular mediated immune response to HIV Gag protein with the recombinant adenoviral particle.
6	25, 26, 44, 45	The claim is directed to a method of generating a cellular mediated immune response to HIV Gag protein with the recombinant adenoviral particle in addition to administering a DNA plasmid vaccine.
7	48-51, 53, 54, 56	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 1) inserted in the parallel orientation of E1.
8	48-51, 53, 54, 56	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 5) inserted in the parallel orientation of E1.
9	48-51, 53, 54, 56	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 7) inserted in the parallel orientation of E1.
10	52	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 1) inserted in the antiparallel orientation of E1.
11	52	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 5) inserted in the antiparallel orientation of E1.
12	52	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 7) inserted in the antiparallel orientation of E1.
13	55	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$

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		and ΔE3, the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 1)
		inserted in E1.
4	55	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 5) inserted in E1.
15	55	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Pol protein (SEQ ID NO: 7) inserted in E1.
16	57-61	The claims are directed to a method of making and harvesting of a recombinant adenoviral particle that contains a gene encoding an HIV Pol protein.
17	62, 65, 66	The claim is directed to a method of generating a cellular mediated immune response to HIV Pol protein with the recombinant adenoviral particle.
18	63, 64	The claim is directed to a method of generating a cellular mediated immune response to HIV Pol protein with the recombinant adenoviral particle in addition to administering a DNA plasmid vaccine.
19	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 9) inserted in the parallel orientation of E1.
20	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 11) inserted in the parallel orientation of E1.
21	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of <u>AE1</u> , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an <u>HIV Nef protein (SEQ ID NO: 13)</u> inserted in the parallel orientation of E1.
22	67-70, 72, 73, 75	The claims are directed to an adenoviral vector that is at least partially deleted of <u>\Delta E1</u> , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an <u>HIV Nef protein (SEQ ID NO: 15)</u> inserted in the parallel orientation of E1.
23	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta EI$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 9) inserted in the antiparallel orientation of E1.
24	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 11) inserted in the antiparallel orientation of E1.
25	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta EI$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 13) inserted in the antiparallel orientation of E1.
26	71	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 15) inserted in the antiparallel orientation of E1.
27	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 9) inserted in E1.
28	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 11) inserted in E1.
29	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$ , the vector contains the cis-acting packaging sequence of the wild type

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· · · ·	1	adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 13) inserted in E1.
30	74	The claim is directed to an adenoviral vector that is at least partially deleted of $\Delta E1$ and $\Delta E3$ , the vector contains the cis-acting packaging sequence of the wild type adenovirus genome, and a gene which encodes an HIV Nef protein (SEQ ID NO: 15) inserted in E1.
31	76-80	The claims are directed to a method of making and harvesting of a recombinant adenoviral particle that contains a gene encoding an HIV Nef protein.
32	81, 84, 85	The claims are directed to a method of generating a cellular mediated immune response to HIV Nef with the recombinant adenoviral particle.
33	82, 83	The claims are directed to a method of generating a cellular mediated immune response to HIV Nef with the recombinant adenoviral particle in addition to administering a DNA plasmid vaccine.
34	86a	The claim is drawn to a multivalent vaccine wherein gag, pol and nef are expressed from three individual vectors.
35	86b, 88, 89	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from one individual vectors.
36	86c, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from two individual vectors, one expressing nef-pol fusion and one expressing gag.
37	86d, 87, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from two individual vectors, one expressing gag-pol fusion and one expressing nef.
38	86e, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from two individual vectors, one expressing nef-gag fusion and one expressing pol.
39	86f, 88	The claims are drawn to a multivalent vaccine wherein gag, pol and nef are expressed from a single vectors as a fusion protein.
40	86g, 88	The claims are drawn to a multivalent vaccine wherein gag and pol are expressed from two individual vectors.
41	86h, 88, 89	The claims are drawn to a multivalent vaccine wherein gag and pol are expressed individually from one vector.
42	86i, 88	The claims are drawn to a multivalent vaccine wherein pol and nef are expressed from two individual vectors.
43	86j, 88, 89	The claims are drawn to a multivalent vaccine wherein pol and nef are expressed from individually from one vector.
44	86k, 88	The claims are drawn to a multivalent vaccine wherein nef and gag are expressed individually from one vector.
45	861, 88, 89	The claims are drawn to a multivalent vaccine wherein nef and gag are expressed individually from one vector.
46	86m, 88	The claims are drawn to a multivalent vaccine wherein gag and pol are expressed as a fusion protein from one vector.
47	86n, 88	The claims are drawn to a multivalent vaccine wherein pol and nef are expressed as a fusion protein from one vector.
48	860, 88	The claims are drawn to a multivalent vaccine wherein nef and gag are expressed as a

The inventions listed as Groups 1-48 do not relate to a single general inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons:

The technical feature linking groups 1-33 appears to be a recombinant adenoviral vector wherein the adenoviral vector is at least partially deleted in E1 but the vector may contain more deletions, the vector contains wild type sequences including packaging signals and a gene encoding a heterologous HIV protein or fragments thereof. Ertl et al. (WO 96/39178) disclose a recombinant adenoviral vector that is deleted in E1 and partially deleted in E3, the remainder of the adenoviral vector contains wild type sequences. The vector additionally contains an insertion of a heterologous protein which includes HIV proteins (see abstract and claims 1 and 5). Therefore, the technical feature linking the inventions of groups 1-45 does not constitute a special technical feature as defined by PCT Rule 13.2, as it does not define a contribution over the prior art.

The special technical feature of the following groups 1-3, 7-15, 19-30 and 34-48 is considered to be the combination of sequences that is disclosed in each group, see individual claim groupings above for the different sequences. The DNA disclosed in each group is made up of a different sequence having a different structure and different function.

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The special technical feature of group 4, 16 and 31 is considered to be a method of producing recombinant adenoviral particles. Each group contains different sequences hence the resulting particles would have different structures and functions associated with the particle.

The special technical feature of group 5, 17 and 32 is considered to be a method of producing a cellular mediated immune response to the heterologous protein encoded by the different adenoviral vectors. Each group contains different sequences a encoding different protein, therefore the resulting immune response will also be different.

The special technical feature of group 6, 18 and 33 is considered to be a method of producing a cellular mediated immune response to the heterologous protein encoded by the different adenoviral vectors in conjunction with immunizing the individual a DNA plasmid vaccine. Each method contains different sequences encoding a different protein, therefore the resulting immune response will also be different.

Accordingly, groups 1-48 are not so linked by the same or corresponding technical feature as to form a single general inventive concept.

#### Continuation of B. FIELDS SEARCHED Item 3:

WEST 2.0, STN-BIOSIS, MEDLINE

adenoviral vector, deletion, HIV, Gag, polyadenylation signal, CMV promoter